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Characterization Of A Putative SIR2 Like Deacetylase And Its Role In SABP2 Dependent Salicylic Acid Mediated Pathways In Plant

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Keywords: Plant defense, Salicylic Acid, SABP2, SBIP-428, SIR2 deacetylase, Acetylation.
ABSTRACT

Characterization of a Putative SIR2 Like Deacetylase and Its Role in SABP2 Dependent Salicylic Acid Mediated Pathways in Plant

by

MD Imdadul Haq

Salicylic Acid Binding Protein2 (SABP2) is an enzyme known to play an important role in the SA mediated pathway. SBIP-428 (SABP2 Interacting Protein-428), an SIR2 like deacetylase, has been found to interact with SABP2. We demonstrate that SBIP-428 functions as a SirTun deacetylase and that SBIP-428 itself is lysine acetylated. Interactions of SBIP-428 with SABP2 also increased the possibility of SABP2 itself being lysine acetylated. Both recombinant purified SABP2 and native partially purified SABP2 displayed no acetylation. In response to TMV infection, expression of SBIP-428 was downregulated at 48 hpi. Additionally, SBIP-428 was upregulated in plants known to accumulate less SA. This evidence suggests that the expression of SBIP-428 is negatively correlated to levels of SA in plants. The AtSRT2 plants exhibit no altered growth phenotype but exhibit higher resistance to bacterial pathogens. Our results indicate that SBIP-428 is an important regulator in the plant defense pathway.
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CHAPTER 1

INTRODUCTION

From archaea to eukaryotic cells, all biotic living organisms use plants as their primary source of food. Some microorganisms, parasitic insects, nematodes, and even parasitic plants use plants to complete their life cycle. Extensive range of organisms attack plants for various reasons. In agriculture, pathogen and herbivores attack can cause serious damage to the quality and yield of crops. Plants have evolved mechanisms to organize and assemble their own defense mechanism to deal with attacks by diverse organisms. In this study, we have tried to understand the defense signaling mechanism of plants.

**Plant Immune System**

Pattern-triggered immunity (PTI) and effector-triggered immunity (ETI) are 2 distinguished models of plant immunity defined by recognition of different types of molecules by plants that are affected by pathogen attack (Jones and Dangl 2006). PTI occurs in plant cell membrane by recognition of microbe-associated molecular patterns (MAMPs) or damage-associated molecular patterns (DAMPs) by plant pattern-recognition receptors (PRR) (Boller and Felix 2009). ETI is triggered upon recognition of pathogen effectors by plant resistance (R) proteins that are normally localized inside the cell (Jones and Dangl 2006).

PTI and ETI have both have strengths and weaknesses; ETI signaling evolved to be a strong response against pathogen effector because of high recognition specificity; however, some pathogen can often escape ETI by eluding such specific recognition. PTI can provide immunity against potential pathogens that are not well adapted but is not very effective against well-adopted pathogens (Tsuda et al. 2009). However, how an effector trigger’s immune response in
plants, coordinates resistance to a broad range of pathogens and their corresponding effector is a question yet to be answered (Chisholm et al. 2006).

Domain Structure of Pattern-Recognition Receptors (PRR) and Resistance (R) Proteins

‘PRR’ and ‘R’ proteins are the 2 major types of proteins involved plant resistance mechanisms. PRRs are associated with or contain interleukin-1 receptor-associated kinase (IRAK) family that is a member of monophyletic group that includes Arabidopsis Flagellin Sensitive2 (FLS2), rice XA21 (Xanthomonas resistance 21), human interleukin-1 receptor-associated kinase (IRKs) and Drosophila Pelle. Pathogen recognition at the surface of cell is carried out by Receptor-Like Proteins (RLPs) and serine-threonine receptor kinases that are known class of PRRs. Receptor kinases XA21 and FLS2 and the RLPs, Cf-9, and XA21D are the best-studied plant PRRs in recent years (Dardick and Ronald, 2006).

Two major classes of protein encoded by resistance (R) genes are; NB-LRR class and eLRR (extracellular Leucine Rich Repeat). They are named after their characteristics nucleotide binding (NB) and leucine rich repeat (LRR), and extracellular leucine rich repeat (eLRR) domains. The length of LRR is usually 20-30 amino acid and the motifs have been identified in proteins with wide range from viruses to eukaryotes. These proteins are involved in various processes from plant development to disease resistance. The subclasses of NB-LRR are coiled-coil (CC) NB-LRR and Toll-interleukin-1 receptor (TIR) NB-LRR. They are divided based on their N-terminal domain. The best characterized member of NB-LRR proteins are RPS2, RPM1, and RPS5. These Arabidopsis R proteins specify resistance to Pseudomonas syringae carrying the bacterial effectors AvrRpt2, AvrRpm1/AvrB, and AvrPphB, respectively (Chisholm et al. 2006). NB-LRR proteins determine the resistance to viral, bacterial, oomycete, and fungal pathogens (Dangl and Jones 2001). eLRR proteins can be further divided according to their
Signaling Pathways and Cross Talk Between JA, ET, and SA.

Jasmonate (JA), ethylene (ET), salicylate (SA), and PAD4, which are termed as ‘signaling sectors’ of a network rather than “pathways”, synthesized with the help of DDE2, EIN2, SID2, and PAD4 genes, respectively. The product of these genes function as hubs that interact with each other. The fragment of bacterial flagellin, flg22 (flg22-PTI) and Pseudomonas syringae effector, AvrRpt2 trigger the PTI and ETI, respectively. The interaction between 4 sectors and flg22-PTI was synergistic, whereas, AvrRpt2-ETI and 4 sectors were compensatory; this difference is not because of the nature of signaling machinery they use but because of how common signaling machinery they use (Tsuda and Katagiri 2010). Major parts of the networks are not different; the speed of the responses affects which immune responses deliver effective immunity (Tsuda et al. 2009). For example, AvrRpm1-ETI responses are markedly faster comparing with AvrRpt2-ETI (Ritter and Dangl 1996; Tao et al. 2003). The signal of pathogen attack branches into multiple signal flows and that multiple signals enter from different points of the common network, which suggests that plants use an integrated signaling network for various modes of immunity but use the common network differently for different modes (Katagiri and Tsuda 2010).

Systemic Acquired Resistance

Systemic acquired resistance (SAR) is a mechanism in which resistance spreads away from the site of primary infection to the uninfected parts of the plant. In this mechanism plants
induce resistance that confers long-lasting defense against a wide spectrum of pathogens
(Durrant and Dong 2004) (Figure 1). Increased level of SA is associated with activation of SAR.
SAR is characterized by coordinate expression of a set of *Pathogenesis-Related (PR)* genes,
many of which encode PR protein with antimicrobial activity (Van Loon et al. 2006). There are
several reports using transgenic or mutant plants to verify the hypothesis that the perception or
production of SA has a key role in SAR? (Loake and Grant 2007; Volt et al. 2009). NPR1, a
regulatory protein, which plays a vital role as transducer of the SA signal, is activated by SA and
acts as coactivator of *PR* gene expression (Dong and Durrant 2004). SA, JA, and ET also play
roles in inducing SAR. There are several reports that suggested that SAR is not activated in
*sgt1b, oopr3, jar, eds8,* and *coi1* *Arabidopsis* mutants that are known JA signaling mutants
(Pieterse et al. 1998; Ton et al. 2002a; Cui et al. 2005; Truman et al. 2007; Attaran et al. 2009).
Verberne et al. (2003) reported that ethylene had a role in SA-dependent SAR induction against
tobacco mosaic virus (TMV). However, the exact role of JA and ET needs to be further studied.
Liu et al. (2011) proposed that interaction of 2 mobile signals: MeSA and a complex of lipid
transfer protein DIR1 and a glycerolipid, control SAR. SAR induction by MeSA is determined
by light period; MeSA and its metabolizing enzymes showed influential effect on SAR during
late in day when plants are expected to receive little light in hours after infection (Liu et al.
2009).

Recently, scientist started to look into long-lasting impact on gene expression and plant
immunity by analyzing epigenic regulatory mechanism, such as chromatin remodeling and DNA
methylation (Bruce et al. 2007; Van Den Burg and Takken 2009; Alvarez et al. 2010).
Figure 1: Activation of Systemic Acquired Resistance in Plant System (Re-drawn with permission from Durrant and Dong 2004). Resistance (R) proteins activate Enhanced Disease Susceptibility 1 (EDS1) and Reactive Oxygen Species (ROS) which activate SA and downstream cascades.

Induced Systemic Resistance

Both pathogenic and nonpathogenic microbes can elevate the disease resistance level in plants. Van loon et al. (1998) first reported that colonization of plant roots by PGPR (plant growth-promoting rhizobacteria) were shown to defend above ground plant tissue against pathogenic microbes. Many nonpathogenic *Pseudomonas* spp and *Bacillus* spp have been documented to be responsible for induced systemic resistance in many plants (Kloepper et al. 2004; Van Loon and Bakker 2006).
Salicylic Acid Synthesis Pathway

SA has been studied for more than 200 years in medicine for humans, but in plants its role is documented only in past ~30 years (Vlot et al. 2009). Wide varieties of role is played by SA in plants, particularly, influencing seed germination, seedling establishment, cell growth, respiration, senescence-associated gene expression, stomata closure, abiotic stress responses, basal thermo-tolerance, nodule formation in legumes, and fruit yield.

Two distinct enzymatic pathways are known to generate SA in plants. Salicylic acid can be produced from chorismate via isochorismate catalyzed by isochorismate synthase (ICS) and isochorismate pyruvate lyase (IPL) (Verberne et al. 2000; Wildermuth et al. 2001; Strawn et al. 2007). On the other hand, SA also can be generated via phenylalanine, cinnamic acid, and benzoate intermediates or coumaric acid (Figure 2). SA glucosyltransferase (SAGT) catalyzes the conversion of SA into SA O-β-glucoside (SAG), or salicyloyl glucose ester (SGE) but how SA is regenerated back from SGA or SGE is still unknown (Dean and Delaney 2008). SA methyltransferase (SAMT) converts SA to MeSA, whereas SABP2/methyl esterase (MES) helps to resynthesize SA from MeSA (Kumar and Klessig 2003).
Figure 2: Synthesis of SA from Phenylalanine or Chorismate in Plant (re-drawn with permission from Volt et al. 2009). SA can be converted from SGE, SAG, MeSA. SABP2 helps in conversion of MeSA to SA.

SA plays extensive role in plant defense signaling against pathogen in local disease resistance mechanism, including death of host cell and defense gene expression, and SAR. PR genes are expressed by SA during resistance response to bacterial, viral, and fungal pathogen infections in plants.

**Salicylic Acid Binding Protein2 (SABP2)**

Salicylic acid binding protein_2 (SABP2), a 29kDa enzyme catalyzes the conversion of MeSA to SA (Kumar and Klessig 2003). Abundance of SABP2 is extremely low in tobacco.
plants but it binds SA with high affinity. Kumar and Klessig (2003) first reported the role of SABP2 in plant innate immunity. The tobacco SABP2 silenced plants are more susceptible to TMV and exhibits down regulation of PR-1 gene expression. In SA signaling pathway, SABP2 plays a significant role in local resistance and SAR. SABP2 likely helps to increase the cytoplasmic SA levels that result in changes in redox potential of cytoplasm. Monomerization of NPR1 oligomers is initiated upon SA mediated changes in redox potential (Beckett and Dorothy 2001). NPR1 monomers migrate to nucleus where TGA class of transcription factors help to induce the expression of SA responsive defense genes resulting in activation of local resistance (Mou et al. 2003). Tripathi et al. (2010) recently reported that SABP2 is required for acibenzolar S-methyl (ASM) (a synthetic analog of SA) mediated activation of SAR in tobacco plants. To better understand the role played by SABP2, a yeast 2-hybrid (Y2H) screen was performed to identify proteins that interact with SABP2.

**SABP2 Interacting Proteins**

Tobacco leaf proteins were used as prey (Y2H library) and full length SABP2 was used as a bait. Several interactors of SABP2 were identified in this screening. One of the interacting proteins identified is SBIP-428. SBIP-428 shows high homology to Silent Information Regulator 2 (SIR2) family proteins.

**Silent Information Regulator2 (SIR2)**

SIR2 proteins, also known as sirtuins, are broadly conserved from bacteria to mammals (Frye 2000). In mammals Sirtuins are divided into 7 groups (SIRT1-SIRT7) (Frye 2000). SIRT1 is the closest homologue of SIR2. Deacetylation of acetylated nuclear histones is well known function of SIR2 that is NAD⁺ dependent (Northa and Verdin 2004). However, some recent studies show that members of SIR2 protein family are also involved in deacetylation of non
histone proteins in other organelles. Enzymatic activity of SIR2 in yeast is required for the association with the regions distal to the nucleation sites (Moazed 2001). Several chemical compounds have been reported to inhibit or activate the deacetylase activity of SIR2. Splitomicin and sirtinol inhibit the NAD$^+$-dependent deacetylase activity of SIR2 in yeast and Arabidopsis (Bedalov et al. 2001; Grozinger et al. 2001). Nicotinamide, a product of SIR2 deacetylation reaction also acts as inhibitor of its activity (Bitterman et al. 2002). Quercetin and piceatannol have been shown to increase the activity of SIR1 proteins (Howitz et al. 2003). Hypoacetylation (decreased acetylation) of histone H4 in the rDNA site was shown to be SIR2-dependent (Guarente and Hekimi 2003). Deletion of SIR2 in yeast, resulted in enhanced acetylation in subtelomeric region, mating type loci, and H3 and H4 histone of rDNA (Buck et al. 2002; Robyr et al. 2002). In humans lack of SIR1 (ortholog of SIR2) showed dependency of embryonic development and muscle differentiation (Cheng et al. 2003; Fulco et al. 2003; Lin and Elledge 2003; McBurney et al. 2003).

*Drosophila* SIR2 (dSIR2) is required for physical interaction with Hairy, a bHLH euchromatic repressor and key regulator of *Drosophila* development (Rosenberg and Parkhurst 2002). dSIR2 is the nuclear protein in adult fly, whereas during embryo stage, prior to nuclear cycle 12, dSIR2 is detected in both nucleus and cytoplasm (Newman et al. 2002; Rosenberg and Parkhurst 2002). Dependency on NAD$^+$ for histone deacetylase activity of SIR2 suggested a link between caloric restriction and SIR2 activity that resulted in extension of life span. Short life span of fly resulted as dSIR2 was deleted (McBurney et al. 2003). Free cytoplasmic NAD$^+$ is a result of less carbon flow through glycolysis under condition of caloric restriction. In circumstances of caloric restriction, NAD$^+$ levels increase that result in SIR2 activation and decrease in the rate of aging. However, Burnett and colleagues (2011) reported that lifespan of
Drosophila and C. elegans increased with dietary restriction independently of dSIR2, questioning the effect of sirtuins on the lifespan of metazoan.

SIR2 in Plants

The functions of SIR2 proteins in plants is not well studied. Recently homolog of SIR2 proteins have been reported in rice and Arabidopsis (Huang et al. 2007; Wang et al. 2010; König et al. 2014). In rice SIR2-like gene (OsSRT1) is needed as safeguard against genome instability and cell damage to ensure plant cell growth. OsSRT1 gene is involved in acetylation and dimethylation of histone H3K9. Knock-down of OsSRT1 resulted in cell death, DNA fragmentation, increased production of H₂O₂ and on the other hand, overexpression showed higher tolerance to oxidative stress (Huang et al. 2007). Wang and colleagues (2010) reported that in Arabidopsis, AtSRT2 acted as a negative regulator of basal defense by suppressing SA biosynthesis. AtSRT2 was down-regulated in response to P. s. pv. tomato DC3000 infection. Expression of PRI was negatively regulated by AtSRT2. HDA19 (histone deacetylase 19), a protein of RPD3/HAD1 superfamily is directly associated with chromatin modification of PRI and PR2 promoter to a repressive state that ensures low basal expression of these genes (Choi et al. 2012). Loss of HDA19 activity increased the level of SA content and expression level of PRI, PR2 (Choi et al. 2012). Zhou et al. (2005) reported that overexpression of HDA19 increased the expression of JA/ET-regulated genes and resistance to fungal pathogen, Alternaria brassicicola, whereas HDA19 knock-down plants were more susceptible. They also reported that HDA19 expression was induced by inoculation with A. brassicicola, wounding, and by the treatment with JA and ET. However, Tian et al. (2005) reported that a histone deacetylase (AtHD1) provides positive and negative control of transcription. AtHD1 gene is involved in ectopic expression of protein synthesis, whereas a negative regulator of SA in leaves and flowers. Microarray analysis
suggested that AtHD1 is involved in regulation of developmental and environmental gene expression (Tian et al. 2005).

Nonhistone Protein Acetylation and Metabolic Enzymes

Several recent studies suggested that beyond histones, a number of other metabolic enzymes in almost all organisms are regulated by acetylation/deacetylation (Frankemier et al. 2011). Acetyl-CoA and NAD+ are most common influencing molecules of protein acetylation that regulates several enzymes activities (Xing and Poirier, 2012). An animal transcription factor, p53, is the other well-known acetylated protein beyond histones (Gu and Roeder 1997). To date, more than 100 other acetylated proteins have been identified in animal, bacteria, and yeast, which include a large number of DNA associated proteins, transcription factors, and nuclear receptors (Kouzarides 2000). Two recent papers have shown that more than 125 different proteins are acetylated in Arabidopsis, which are involved in wide range of metabolic pathways (Frankemier et al. 2011; Wu et al. 2011).
Hypotheses

Hypothesis I: SBIP-428, a homologue of Silent Information Regulator 2 (SIR2), is a NAD$^+$ dependent deacetylase.

Hypothesis II: SABP2 is acetylated and is regulated by deacetylase activity of SBIP-428.

Hypothesis III: SBIP-428 has an effect on SA mediated defense mechanism in plants.
CHAPTER 2

MATERIALS AND METHODS

Materials

Plant Materials

Three transgenic lines of tobacco plants *Nicotiana tabacum* cv. *Xanthi nc NN* (XNN), NahG (expressing *nahG* gene, that encodes SA hydrosylase that converts SA to catechol), 1-2 (SABP2 silenced) were used for this study. Six mutant lines of *Arabidopsis thaliana* (CS370961, SALK_139443, SALK_131994C, SALK_149295C, SALK_035541, and CS877409) and a wild type (Col-0) also were used for this study. Seeds were grown in autoclaved (for 20min) soil containing peat moss (Fafard Canadian growing mix F-15, Agawam, MA). Tobacco seedlings were transferred to 4 X 4 inch flats after 14 days, 2 in one flats. After 30 days, young plants were transfer to an 8 inch pot. Fertilizer was added to the plants after 3 days of transfer to the pot. Six to 8 weeks old plants were used to perform the experiments. *Arabidopsis* seeds were sown in autoclaved soil with fertilizer. Seeds were kept in the cold dark room for 3 days and then transfer to the growth chamber. The growth chamber (PGW 36, Conviron, Canada) was set at 16-h day cycle with temperature at 22°C.

Chemicals and Reagents

Sodium dodecyl sulfate (SDS), ß-mercaptoethanol (ß-ME), tetramethylethlenediamine (TEMED), ammonium persulfate (APS), bovine serum albumin (BSA), bovine thrombin, coomassie brilliant blue R-250, coomassie brilliant blue G-250, ponceau-S, ethylenediaminetetraacetic acid (EDTA), TRIS base, phenylmethylsulfonyl fluoride (PMSF), glycine, glycerol, methanol, imidazole, Tween-20, Triton X-100, N,N-Bis(2-
hydroxyethyl)glycine (Bicine), magnesium chloride (MgCl$_2$), sodium chloride (NaCl), sodium phosphate monobasic (NaH$_2$PO$_4$), sodium phosphate dibasic (Na$_2$HPO$_4$), benzamidine-HCl, ammonium sulfate ((NH$_4$)$_2$SO$_4$), sodium citrate (Na$_2$H$_5$O$_7$C$_6$), and all other standard chemicals were purchased from Fisher Scientific, Pittsburgh, PA. Polyvinylpolypyrrolidone (PVPP) was purchased from Acros Organics, Audubon Park, NJ. The Mini Protean 3 cell, 30% acrylamide, Bradford’s reagent, prestained low molecular weight marker, 10x SDS loading buffer, SDS dye, and Mini trans blot system were purchased from Bio-Rad, Hercules, CA. Polyvinylidene fluoride (PVDF) membranes were purchased from Millipore, Billerica, MA. Mono Q column was purchased from GE Healthcare, Piscataway, NJ. Bicinchorinic acid (BCA) protein assay and Pierce ECL western blotting substrate were purchased from Thermo Scientific, Rockford, IL. Kodak developer and fixer replenisher were purchased from Sigma-Aldrich, St. Louis, MO. 6xHis-tag SBIP-428 protein was expressed in Gateway system. Rabbit polyclonal SABP2 antibodies and monoclonal anti-rabbit, IgG $\gamma$ chain specific secondary antibodies conjugated to HRPO and tobacco mosaic virus (TMV) were available in-house. Oligo dT-20, Taq DNA polymerase (Invitrogen, CA), dithiothreitol (DTT), DNA ladder (New England Biolabs), MMLV reverse transcriptase, RNase free DNase, recombinant RNAsin (Promega), and gel loading dye (Bio-Rad).

**Other Materials**

One ml syringes (BD syringes, NJ), pestle grinder (Fisher Scientific), cheese cloth and miracloth (Fischer Scientific), Spectrophotometer, high speed centrifuge (Beckman, model J2-21 or Sorvall RC5B), SYNERGY HT Multi-Mode Microplate Reader (Biotek), and AKTA purifier 10 (GE Healthcare) system were used for this research.
Methods

Biochemical Characterization (Hypothesis I)

Bioinformatics Analysis

Sequence Alignments and Database Analysis. The DNA sequence and corresponding amino acid sequence of SBIP-428 were analyzed using BLAST (Basic Local Alignment Search Tool) (Altschul et. al. 1990) at NCBI (http://blast.ncbi.nlm.nih.gov/Blast.cgi) and ExPASy Bioinformatics Resource Portal (http://www.expasy.org/tools/). To compare it with known tobacco genes, the SBIP428 nucleotide sequence was used to search the tobacco Unigene in solanaceae database (http://solgenomics.net/tools/blast/index.pl). Full length protein sequences of *Ricinus communis*, *Populus*, *Arabidopsis thaliana*, and *Oryza sativa* similar to SBIP428 were compared using Multi-sequence alignments of amino acid (ClustalW2; http://www.ebi.ac.uk/Tools/msa/clustalw2/).

3D Protein Structure Model of SBIP-428. (PS)²: Protein Structure Prediction Server (http://ps2.life.nctu.edu.tw/) was used to generate the 3D protein structure of SBIP-428. A human Sir2 was used as template to generate the 3D structure models.

Subcellular Localization Prediction of SBIP-428. Protein Prowler (http://bioinf.scmb.uq.edu.au/pprowler_webapp_1-2/), MultiLoc (http://abi.inf.uni-tuebingen.de/Services/MultiLoc/), and BaCelLo (http://gpcr.biocomp.unibo.it/bacello/pred.htm) prediction tools were used to predict the subcellular localization of SBIP-428.

Expression and Purification of SBIP-428

Cloning of SBIP-428. The cloning of full length SBIP-428 in pDONR221 was previously performed (Kumar et al., unpublished). Briefly, the full length coding region of SBIP-428 was
amplified by RT-PCR and cloned into entry plasmid vector pDONR221 (Gateway, Invitrogen). Obtained cDNA was cloned, sequenced, and analyzed. Once a full length SBIP-428 was obtained and verified, it was used for recombinant protein expression. To express recombinant proteins, the full length SBIP-428 was sub-cloned into pDEST17 (Gateway; Invitrogen) with 6XHis tag on its N-terminus. Following verification of sequences, the recombinant SBIP-428 in pDEST 17 was transformed into a suitable *E. coli* host (BL21 (DE3) pLysS and BL21-CodonPlus (DE3)-RIL) for protein expression.

**Purification of SBIP-428 Using Ni-NTA Chromatography.** Recombinant his-tagged SBIP-428 was first purified by Ni-NTA column chromatography. A modified purification protocol described by Forouhar et al. (2005) was used. The *E. coli* BL21 pLysS cells were grown in the LB agar containing 100 µg/ml ampicillin at 37°C for overnight. The overnight grown single colony was inoculated to 3ml of liquid LB broth (containing 100 µg/ml of ampicillin). The culture was grown for ~12 hour then diluted 100 times in fresh 50 ml LB broth with antibiotics. The overnight grown culture was then diluted to 1.5 L of fresh LB medium (containing antibiotic). Bacterial cultures were grown at 37°C in a shaker maintained at 250 rpm until it reached 0.6-0.7 OD (optical density). At OD= 0.6-0.7, protein expression was induced by adding 0.01 mM of isopropyl β-D-1 thiogalactopyranoside (IPTG) to the culture and was incubated at 17°C in a shaker maintained at 250 rpm for overnight. Overnight culture was centrifuged at 8000 rpm at 4°C for 10 minutes to pellet bacteria. Bacterial pellet was then suspended in 1X Ni-NTA buffer (50 mM Na₂HPO₄, 300 mM NaCl, and 10 mM Imidazole, pH 8.0) containing 0.1 mM protease inhibitors (phenylmethylsulfonyl fluride (PMSF); 100mM stock in isopropanol) and then lysed using French-press (Thermo Electron Corporation; cell type 20 K, max gauge 1280). Cell lysate was centrifuged at 14000 rpm for 15 min at 4°C to separate soluble (supernatant) and
insoluble (pellet) proteins. Soluble proteins in supernatant were purified by using metal affinity chromatography (Ni-NTA). Ni-NTA resin pre-equilibrated in 1X Ni-NTA buffer (50 mM Na₂HPO₄, 300 mM NaCl, and 10 mM Imidazole, pH 8.0) was added to soluble protein and incubated at 4ºC with gentle shaking for overnight. Flow-through containing unbound proteins was collected and loosely bound proteins was washed using 1x Ni-NTA buffer. Bound proteins were eluted using 250 mM of imidazole in 50 mM Na₂HPO₄, 300 mM NaCl, pH 8.0. Purification of SBIP-428 was analyzed by SDS-polyacrylamide gel electrophoresis and western blot was performed using monoclonal Anti-polyHistidine antibody to confirm the presence of SBIP-428.

Purification Using Mono-Q. Fractions containing SBIP-428 from the Ni-NTA chromatography were further purified on a mono-Q anion exchange column. Fractions containing SBIP-428 from Ni-NTA were pooled and desalted using buffer (10 mM Bicine, pH 8.0) in a Hitrap Desalting Column (2x5 ml) (GE Healthcare). Desalted fractions were applied to Mono-Q column (Mono-Q 5/50 GL, GE Healthcare). The bound proteins were eluted with a linear gradient of 0-300 mM ammonium sulfate in buffer (10 mM Bicine, pH 8.0). The presence of SBIP-428 was determined by western blot using monoclonal Anti-polyHistidine antibody (1:3000 dilution in blocking buffer; 1 % BSA, 3 % nonfat milk in 1xPBS), and a 12.5 % SDS-PAGE was run to confirm purification of SBIP-428.

Dialysis of SBIP-428. Mono Q purified proteins (~4 ml) were pooled together in a dialysis bag (Molecular Weight cutoffs: 12000-14000 Da). Samples were dialyzed against deacetylase buffer (50 mM Hepes pH 7.6, 350 mM NaCl, and 20 % glycerol) at 4ºC with gentle stirring. Buffer was changed every 8 hours 3 times. Dialyzed proteins were aliquoted (100µl) and stored at -80ºC until further use.
Deacetylase Activity of SBIP-428

Deacetylase Enzyme Assay Using Total Leaf Protein as Substrate. A deacetylase enzyme assay was performed as described by Frankemier et al. (2011) with minor modifications. Total acetylated proteins from the tobacco leaves were used as substrate for the enzyme assay. Plant proteins are naturally acetylated and could potentially be used for deacetylation assay. Total leaf proteins were extracted as described below using direct extraction buffer (DEB; 100 mM Tris-HCl pH 8.0, 1.5 M β-mercaptoethanol, 15 % glycerol, 5 mM NaF, 1 mM Na3VO4, and 2 mM EDTA). Proteins were desalted using a spin column prior to the enzyme assay. The SBIP-428 enzyme was purified using mono-Q ion exchange chromatography described as above.

For deacetylase activity assay of recombinant SBIP-428, 12 µg of *N. tabacum* (XNN) leaf protein (naturally acetylated) were incubated with 18 µg, 12 µg, or 8 µg of SBIP-428 for 3 h at 37°C. As a negative control (without SBIP-428), deacetylase buffer (50 mM Hepes, pH 7.6, 350 mM NaCl, 20 % Glycerol) was used instead of SBIP-428. Proteins treated with the enzyme SBIP-428 were then subjected to Western blot analysis using anti Acetylated Lysine Antibody. Experiment details are presented in Table 1 below-

### Table 1: Details of SBIP-428 Enzyme Assay Work Flow

<table>
<thead>
<tr>
<th></th>
<th>Proteins (substrate)</th>
<th>SBIP-428 (enzyme)</th>
<th>Buffer (5X)</th>
<th>Water</th>
<th>Total Volume</th>
<th>Dye (X)</th>
<th>Loading volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>8µl XNN (12 µg)</td>
<td>11.21 µl (12 µg)</td>
<td>6 µl</td>
<td>4.79 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>2</td>
<td>8 µl XNN (12 µg)</td>
<td>11.21 µl (12 µg)</td>
<td>6 µl</td>
<td>12.79 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>3</td>
<td>8 µl XNN (12 µg)</td>
<td>16.82 µl (18 µg)</td>
<td>6 µl</td>
<td>0 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>4</td>
<td>8µl XNN (12 µg)</td>
<td>16.82 µl (18 µg)</td>
<td>6 µl</td>
<td>7.18 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>5</td>
<td>8µl XNN (12 µg)</td>
<td>5.16 µl (8 µg)</td>
<td>6 µl</td>
<td>10.39 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>6</td>
<td>8µl XNN (12 µg)</td>
<td>5.16 µl (8 µg)</td>
<td>6 µl</td>
<td>12 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>7</td>
<td>8µl XNN (12 µg)</td>
<td>5.16 µl (8 µg)</td>
<td>6 µl</td>
<td>16 µl</td>
<td>30</td>
<td>5 µl (6X)</td>
<td>33 µl</td>
</tr>
<tr>
<td>8</td>
<td>8µl XNN (12 µg)</td>
<td>16.82 µl (18 µg)</td>
<td>6 µl</td>
<td>5 µl (6X)</td>
<td>21 µl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>8µl XNN (12 µg)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>5 µl (6X)</td>
<td>12 µl</td>
</tr>
</tbody>
</table>

XNN; Total leaf protein from *N. tabacum*
SBIP-428 enzyme; Ni-NTA and mono Q purified protein.

**Deacetylase Enzyme Assay Using SIRT GloTM Assay Kit.** Mono-Q purified SBIP-428 was used to determine the enzyme activity using SIRT-Glo™ assay kit (Promega). Protein was expressed, purified, and quantified for this assay. SRT2 deacetylase enzyme activity was determined as described in the manufacturer’s protocol. Briefly, SIRT-Glo™ buffer and substrate were equilibrated at room temperature and developer reagent was thawed at room temperature. SIRT-Glo™ buffer (10 ml) was added to SIRT-Glo™ substrate and then mixed gently to form SIRT-Glo™ substrate solution. Developer reagent (10 µl) was added to SIRT-Glo™ substrate solution to form SIRT-Glo™ reagent. SBIP-428 was serially diluted (1 µg to 8 µg) in total 100 µl of SIRT-Glo™ buffer and dispensed in a 96 well plate along with 100 µl of SIRT-Glo™ substrate solution. The final total reaction volume was 200 µl. Three replication of each concentration were used for this enzyme activity. SIRT-Glo™ substrate solution (100 µl) was added with 100 µl of SIRT-Glo™ buffer as negative control for this experiment. Reaction mixture was mixed briefly (~15 sec) at room temperature (RT) using the orbital shaker at 500-700 rpm and incubated at RT for 15 min. Luminescence measurements was taken at steady-state signal.

**Deacetylase Enzyme Assay Using HDAC GloTM Assay Kit.** Mono-Q purified SBIP-428 was used to determine the enzyme activity using HDAC-Glo™ assay kit (Promega). Histone deacetylase activity was determined as described in the manufacturer’s protocol. SBIP-428 was serially diluted (1 µg to 8 µg) in total 100 µl of HDAC-Glo™ buffer and dispensed in 96 well plate along with 100 µl of HDAC-Glo™ substrate solution. The final total reaction volume was 200µl. Reaction mixture was mixed briefly at room temperature using the orbital shaker at 500-700 rpm and further incubated at room temperature for 15min. Luminescence measurements was
detected at signal steady-state. Three replication of each concentration were used for this enzyme activity. HDAC-Glo™ substrate solution (100 µl) was added with 100 µl of HDAC-Glo™ buffer as negative control for this experiment.

**Lysine Acetylation of SBIP-428.**

Purified SBIP-428 was analyzed for the potential of lysine acetylation. A Western blot was performed to detect the lysine acetylation of SBIP-428.

**SABP2 Acetylation (Hypothesis II)**

**Purification of Recombinant SABP2 Expressed in E. coli**

A modified protocol was used for SABP2 purification described by Forouhar et al. (2005). *E. coli* cells (MGK cells) containing pET28-SABP2 was grown in the LB agar containing ampicillin 100 µg/ml and kanamycin 10 µg/ml for overnight. The overnight grown single colony was inoculated to 3ml of liquid LB medium (containing 100 µg/ml of ampicillin and 10 µg/ml kanamycin). The culture was grown for 12 hours then diluted 100 times in fresh 200ml LB medium containing antibiotics. Bacterial culture was grown at 37°C in a shaker maintained at 250rpm until reach 0.6-0.7 OD. Once desired OD reaches at 0.6-0.7, protein expression was induced by adding 0.01 mM of isopropyl β-D-1 thiogalactopyranoside (IPTG) to the culture and was incubated at 17°C in a shaker maintained at 250 rpm for overnight.

Overnight culture was centrifuged at 8000 rpm for 10 minutes at 4°C to pellet bacteria. Bacterial pellet was then suspended in 1X Ni-NTA buffer (50 mM Na2HPO4, 300 mM NaCl, and 10 mM Imidazole, pH 8.0) containing protease inhibitors and then lysed using French-press. Cell lysate was centrifuged at 14000 rpm for 15min at 4°C. Supernatant containing soluble proteins were purified by using metal affinity chromatography using Ni-NTA (Qiagen). Ni-NTA resin (~4 ml)
was equilibrated in 1x Ni-NTA binding buffer by washing at least 4 times. Soluble protein extract was added to the Ni-NTA resin and incubated with gentle shaking for overnight at 4°C. Next morning, Ni-NTA resin with soluble protein extract was brought to room temperature and poured into an empty column. Resin was allowed to settle down before opening stop valve. Flow-through containing unbound proteins was collected and loosely bound proteins were washed using wash buffer (50 mM Na$_2$HPO$_4$, 300 mM NaCl, and 10 mM Imidazole, pH 8.0). Bound proteins were eluted using 250 mM of imidazole in 1x Ni-NTA buffer. Purification of 6xHis tagged SABP2 was analyzed by SDS-polyacrylamide gel electrophoresis and western blot was performed using SABP2 polyclonal antibody to confirm the presence of SABP2. SDS-PAGE and Western blot protocol described below.

**Purification of Native SABP2 from Tobacco**

Native SABP2 was purified from WT (wild type) tobacco *Nicotiana tabacum* cv. Xanth nc NN (XNN). Fully grown leaves from 6 to 8 week old plants were used for this experiment. Leaves were harvested and then kept frozen in liquid nitrogen. Frozen tissue were ground using a pestle and mortar. Three volumes of protein extraction buffer (20 mM sodium citrate, 5 mM MgSO$_4$, 1 mM EDTA, 14 mM β-mercaptoethanol, 0.1 mM phenylmethylsulfonylflouride (PMSF), 1 mM benzamidine-HCl, and pH-6.3) were added to homogenized tissue sample. All subsequent steps were carried out at 4°C. The homogenate was filtered through 1 layer of miracloth (Calbiochem) and 4 layers of cheesecloth (Electron Microscopy Science). The filtered extract was centrifuged at 10,000 rpm for 20min at 4°C (GS3 rotor, Sorvall). Supernatant were used for further ammonium sulfate fractionation.

**Ammonium Sulfate Fractionation.** To obtain the 0-50% ammonium sulfate precipitated proteins, 50% ammonium sulfate (313g per liter) was slowly added and stirred into the
supernatant (soluble leaf extract). The saturated supernatant was stirred on ice for additional 20 min. After 20 min, the solution was centrifuged at 10,000 rpm for 20 min at 4°C (GS3 rotor, Sorvall). The pellet (containing 0-50% precipitated proteins) was stored at -20°C until use. Supernatant containing soluble proteins were precipitated further with 75% ammonium sulfate (176g per liter) by adding powdered ammonium sulfate. Incubation on ice and centrifugation was done as described above. The pellet was collected as 50-75% precipitated proteins and stored at -20°C for future use. The presence of SABP2 and acetylation of SABP2 was determined by Western blot using SABP2 polyclonal antibody and Anti-Acetyl Lysine Antibody, respectively.

**Purification of Native SABP2 form Tobacco upon TMV Infection**

Six-week-old wild type tobacco plant was infected with tobacco mosaic virus (TMV) with a $10^{-3}$ dilution of 45µg/ml TMV preparation. Carborundum was dusted evenly on the upper surface of the leaves, and TMV diluted in 0.05 M sodium phosphate buffer was gently rubbed on with a piece of cheesecloth. TMV infected plant was kept separated from uninfected plants and exposed with regular water and light conditions (16h day). SABP2 was purified in the same manner that SABP2 from native tobacco was. SDS-PAGE (12.5%) was done to resolve the 50-75% precipitated proteins and a Western blot was performed using polyclonal SABP2 antibodies to detect native SABP2. The Western Blot was performed with the purified protein with Anti-Acetyl Lysine Antibody to determine the acetylation of native SABP2 upon TMV infection.
SBIP-428 is SA Mediate Defense Pathway (Hypothesis III)

Effect of SA on the Deacetylase Activity in Tobacco Transgenic Plants

Total protein was extracted from tobacco leaves using direct extraction buffer (DEB; 100 mM Tris-HCl pH 8.0, 1.5 M β-mercaptoethanol, 15 % glycerol, 5 mM NaF, 1 mM Na$_3$VO$_4$, and 2 mM EDTA). Plant leaves were ground in 3 times of DEB buffer using a pestle and mortar with acid washed sea sand (Fisher Scientific). Extract was then incubated in boiling water bath for 5 minutes and centrifuged for 10 minutes at 13000 rpm at room temperature. The supernatant was collected as protein sample and quantified using Bradford reagent (Bio-Rad). The Western Blot was performed with the purified protein with Anti-Acetyl Lysine Antibody to determine the effect of SA on deacetylase activity.

Expression of SBIP-428 in Tobacco Transgenic Plants

To determine the effect of SA on SBIP-428 expression in *Nicotiana tabacum* cv. Xanthi nc (NN), NahG (express *nahG* gene, that encodes SA hydrosylase which convert SA to catechol), 1-2 (*SABP2* silenced) were used. Three leaf disc samples were collected from each plant using a cork borer (size 4).

Expression of SBIP-428 in Tobacco upon TMV Infection

Six-week-old wild type tobacco plant *Nicotiana tabacum* cv. Xanthi nc (NN) were infected with tobacco mosaic virus (TMV) with a 10$^{-3}$ dilution of 45µg/ml viral concentration. Carborundum was dusted evenly to the upper surface of the leaves, and TMV diluted in phosphate buffer was gently rubbed on with a piece of cheesecloth. TMV infected plant was kept separate from uninfected plants. Plants were kept under light (150 µMm$^{-2}$s$^{-1}$) and day conditions at ~22°C. Three leaf discs were collected at 0, 24, 48, and 72 hpi (hours post inoculation) from
the inoculated leaves of XNN plants. Samples were frozen in liquid nitrogen and stored at -80°C until ready for total RNA isolation.

**Complementation Experiment**

**SIR2 Mutants of Arabidopsis thaliana.** Six different T-DNA insertion lines were obtained from TAIR (arbidopsis.org). The insertion lines are SALK_131994C, SALK_139443, SALK_149295C, SALK_035541, CS370961, and CS877409. Details of T-DNA insertion lines are presented in Table 2 below.

**Table 2: Details of Arabidopsis SRT2 Mutant**

<table>
<thead>
<tr>
<th>Seed Lines</th>
<th>Background</th>
<th>Clone Name</th>
<th>Insertion site</th>
<th>Gene names</th>
<th>Locus</th>
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<td>CS370961</td>
<td>Col-0</td>
<td>pAC161</td>
<td>Intron</td>
<td>Histone deacetylase</td>
<td>AT4G38160</td>
</tr>
<tr>
<td>SALK_13944</td>
<td>Col-0</td>
<td>pROk2</td>
<td>Exon</td>
<td>Histone deacetylase</td>
<td>AT4G38160</td>
</tr>
<tr>
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<td>Intron</td>
<td>SIRTUIN2</td>
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<td>Col-0</td>
<td>pDAP101</td>
<td>Intron</td>
<td>SIRTUIN2</td>
<td>AT5G09230</td>
</tr>
</tbody>
</table>

**Confirmation of T-DNA Insertion.** T-DNA insertion confirmation experiment was carried out following the protocol of the TAIR website. The brief details are as follows: 2 seeds from each seed line were sown in a 3x3 inch flat with autoclaved soil and kept for 3 days in a dark room at 4°C. Seeds were then transferred to a growth chamber with 16h day light (7000 lux). Leaf samples were collected from each mutant line to obtain the DNA and analyze T-DNA insertion.

**DNA Extraction Protocol.** Three to 4 leaf disks (about 3-5mg) were collected in a 1.5 ml micro-centrifuge tube and weighed. 200µl of Extraction buffer {10 fold dilution of Edward’s
solution (200 mM Tris-HCl, pH 7.5, 250 mM NaCl, 25 mM EDTA, 0.5% SDS) in TE buffer (10 mM Tris-HCl, pH 8.0)) was added and homogenized using a hand held tissue grinder. Samples were centrifuged at 14000rpm for 5min at room temperature. Supernatant was collected as a DNA sample. Collected extract (1 µl) was used in total 20 µl of PCR reaction to analyze T-DNA insertion. Primers details of the seed lines are in the Table 3 below.

Table 3: Primer Details of *Arabidopsis SRT2* Mutants

<table>
<thead>
<tr>
<th>Name of Primers</th>
<th>Sequences</th>
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<tr>
<td>CS370961-RP</td>
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<td>CS877409-RP</td>
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<tr>
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<td>ATGATCCGGACATTGTGCTAG</td>
</tr>
<tr>
<td>SALK_131994C-RP</td>
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<tr>
<td>SALK_149295C-LP</td>
<td>CGCAGAGAGAGACAAAAATCG</td>
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<td>SALK_149295C-RP</td>
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<td>SALK_035541-LP</td>
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<td>SALK_035541-RP</td>
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<tr>
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</tr>
</tbody>
</table>

**Confirmation of Mutant.** RT-PCR was used to determine the SIR2 mutant out of these *Arabidopsis* seed lines. Primers were made using the Primer3 software (http://www.bioinformatics.nl/cgi-bin/primer3plus/primer3plus.cgi/) to analyze the mutant and listed in Table 4 below.
<table>
<thead>
<tr>
<th>Gene</th>
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<tr>
<td>Histone Deacetylase</td>
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<td>GGTACACTTGAGCATGCTG</td>
<td>265</td>
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</tbody>
</table>

Growth Phenotype Analysis of Mutant

*Arabidopsis SRT2* mutant (SALK_131994C) and *Arabidopsis* wild type (Col-0) were used to analyze the growth phenotype. Fifty seeds were grown from each line to compare the growth phenotype of mutant. Germination rate, number of leaves, leaves size, number of flower, height of flower bolds were measured to compare the growth phenotype of SRT2 mutant and Col-0. Seeds were sown in autoclaved soil and then kept 4°C in dark and for 3 days before transfer to growth chamber with 700 to 800 µE/m²/sec light at 16 h day period. Germination rate was determined after 7 days of sowing seeds. The number of leaves in each plant was counted, and the length of the leaves was measured after 21 days of germination. Number of flower and height of flower bold were measured between 30 to 35 days.

Bacterial Growth Assay

A recently developed flood-inoculation technique with minor modification described by Ishiga et al. (2011) was used for bacterial pathogenesis assay. Briefly, Arabidopsis seeds were surface sterilized by incubating in 70% ethanol for 5min in a microcentrifuge tube followed by 20% (v/v) commercial bleach containing 0.1% Tween 20. Seeds were washed at least 5 times.
with sterile milli-Q water. Five seeds were then germinated in half strength MS medium containing Gamborg vitamins solidified with 0.8% Phyto-agar in Petri plates (100 mm X 15 mm). Plates were incubated overnight to remove extra moisture. Seeds were transferred to the MS plates and kept in dark for 2 days at 4ºC. Plates with seeds were then transferred to tissue culture room with light intensity of 150-200 µE/m²/sec at 24ºC and a 16 hr light period. Two weeks postgerminated seedlings were used for pathogen assay.

*Pseudomonas syringae* pv. tomato DC3000, and *Pst* DC3000 *AvrRpt2* stains were used for pathogen assay. Bacterial stains were grown in King’s B medium containing rifampicin (50 µg/ml) and kanamycin (25 µg/ml) for overnight. Bacteria were suspended in sterile milli-Q water and OD was measured at 600 nm. Two different concentrations (1 x 10⁸ and 5 x 10⁶ CFU) were used for this experiment. Bacterial suspension (40 ml) was added to 2 weeks old postgerminated seedlings and kept in room temperature for 2-3 min. Bacterial suspension was removed by decantation and then plates were sealed with 3 M Micropore 2.5 cm surgical tape. Plates were transferred in growth chamber with a light intensity of 150-200 µE/m²/sec at 24ºC with 16h day period. Internal bacterial population was measured at 4dpi. Leaves were collected weight was recorded for each sample. Then leaves were surface sterilized with 3% H₂O₂ for 3min. Samples were then washed with sterile milli-Q water at least 4 times. Washed samples were homogenized in sterile mili-Q water and diluted samples were plated onto King’s B medium containing rifampicin (100 µg/ml) and kanamycin (50 µg/ml). Bacterial colony forming units were counted after 2 days using proper diluted samples.
Protocols for RT-PCR (Reverse Transcriptase Polymerase Chain Reaction) and Western Blot

Isolation of Total RNA for Leaves

Leaf discs were ground using a pestle in liquid nitrogen. To the powdered leaf, 1ml of Tri-reagent (Sigma) was added, mixed gently, and incubated for 5 min at room temperature. To the extract, 200 µl of chloroform was added and mixed by inverting several times. Samples were incubated at room temperature for 3 min. The sample was then centrifuged at 12000 x g for 15 min at 4ºC. The aqueous phase was transferred to a new tube, and 0.5ml isopropanol was added. The samples were mixed by inverting several times and incubated for 10min at room temperature. The sample was then centrifuged at 12000 x g for 10 min at 4ºC. The supernatant was discarded, and the pellet was collected for further processing. Ice cold 75% ethanol (1ml) was added to the pellet. The sample was gently mixed and centrifuged at 7500 x g for 5 min at 4ºC. The supernatant was discarded, and the pellet was air dried for 10-20min. The pellet was then suspended in 43 µl of 0.1% depc treated water. For DNase treatment, 5 µl of DNAse buffer and 2 µl of DNAse (Promega) was added to the resuspended RNA and incubated at 37ºC for 20 min. To remove DNase, the sample was treated with TRI-reagent, chloroform, isopropanol, and chloroform with half of the volume as described above. The pellet was collected after the last step and air-dried for 10-20 min. The air-dried sample was suspended in depc treated water and heated at 55ºC for 5-10 min to make sure the pellet was well dissolved in the depc treated water. RNA concentration (ng/µl) was measured by using nanodrop spectrophotometer (NanoDrop Technologies ND-1000) at 260 nm. The concentration was maintained at 1µg/8µl for cDNA synthesis.

cDNA Synthesis

Reverse transcriptase enzyme (Promega) was used to synthesize first strand
complementary DNA (cDNA) from the isolated total RNA. In the first step, 2 μl of oligo-dT (0.5 μg/ml) was added to 8 μl (1 μg) of total RNA and incubated at 75°C for 10 min in the thermocycler (Effendorf). The sample was then cooled down to 4°C and a 10 μl mix (1 μl reverse transcriptase (RT) (MMLV), 4 μl 5X RT buffer, 1 μl RNAsin (RNAase inhibitor), 1 μl 10 mM DNTP, and 3 μl DEPC treated water) was added to the sample. The sample was vortexed briefly and incubated at 42°C for 60 min and then at 70°C for 10 min in the thermocycler. The sample was stored at -20°C for further analysis.

**Reverse Transcriptase-Polymerase Chain Reaction (RT-PCR)**

The cDNA sample was used to analyze gene expressions. Total 10 μl of PCR mix was used to amplify each gene. The PCR mix contained 6 μl depc treated water, 1 μl 10 X dNTP, 1 μl 10 X Taq polymerase buffer, 0.2 μl Taq polymerase, 0.4 μl of 10 μM forward primer, 0.4 μl 10 μM reverse primer, and 1 μl of cDNA sample.

**Agarose Gel Electrophoresis**

Agarose gel electrophoresis was used to analyze the genes using the PCR product. Agarose gel (0.8% to 1.2%) was used according to gene size. Agarose was added to the 1xTAE buffer (40 mM Tris base, 40 mM Acetic Acid, 1 mM ETDA) then heated, and ethidium bromide (10 mg/ml) was added (1μl/20ml) before pouring into a gel tray.

**SDS-PAGE**

Laemmli’s protocol (Laemmli, 1970) was followed to perform SDS-PAGE. Unless stated otherwise, each sample was mixed with equal volume 2X SDS sample buffer containing β-ME, boiled for 5 minutes, and centrifuged at 13000 rpm for 10 minutes at room temperature. Protein separation on SDS-PAGE was performed at 15 mA for one gel or 25 mA for 2 gel at 200
V for 80 to 90 minutes. The buffer composition used is described in the Appendix B.

Western Blot Analysis for SABP2 and His-Tagged Protein

The Western analysis was performed using standard protocol after the electrophoresis gels were incubated in transfer buffer (25 mM Tris, 192 mM Glycine, 10% methanol) for a few minutes (Towbin et al. 1979). Prior to transferring proteins from gel to PVDF membrane, the PVDF membranes were soaked in 100% methanol for 15 seconds following a rinse with milliQ water for 2 minutes and incubated in transfer buffer for 10 minutes. The gel and PVDF membranes were sandwiched between presoaked sponges and Whatman 3mm Chr papers, protein transfers were carried out at a constant 100 V for 1 hour at 4ºC. After transfer, the membranes were placed in 100% methanol for 10 seconds and were let dry on Whatman paper. Membranes were soaked again in 100% methanol again for 10 seconds and then stained with Ponceau-S and photographed to verify the transfer of proteins. Membranes were washed with PBS buffer until the stains were gone and then probed with rabbit polyclonal SABP2 primary antibody (1:1000) or mouse monoclonal anti PolyHistidine primary antibody (1:3000) in 5ml of blocking buffer (3% BSA, 1% milk in 1 x PBS) overnight at 4ºC on a shaker. The blots were washed 3 times for 5 minutes each with 1x PBS, 1x PBST with 3% tween 20, and 1x PBS sequentially. The blot was then probed in either antirabbit secondary antibody or antimouse IgG peroxidase secondary antibody diluted at 1:10000 for 1 hour at room temperature and washed sequentially as described earlier. The signals on membranes were developed with ECL substrate (Thermo Scientific) for 1 minute, and protein expression was analyzed by using X-ray films as described by the manufacturer.

Western Blot Analysis for Lysine Acetylated Proteins

Proteins were transferred from gel to PVDF membrane as described above. PVDF
membrane was then subjected to a blocking buffer for 30 minutes at room temperature. PVDF membrane was then washed with PBS and PBST (PBS with 0.5% Tween 20) 2 times each. The blot was then probed in Anti-lysine Antibody diluted in PBST for 16 to 18 hours at 4°C. The blot was then thrice washed with PBST for 5 minutes. Secondary antibody (anti-rabbit) was added to the blot and kept for 2 hours at room temperature. The signals on membranes were developed with ECL substrate (Thermo Scientific) for 5 minutes, and protein expression was analyzed by using X-ray films as described by the manufacturer.

**Spin Column**

Syringe (1 ml) was used as column. Fiber glass was added to the bottom of 1 ml syringe and then filled with Sephadex G 25 in citrate buffer (20 mM sodium citrate pH 6.5, 1 mM EDTA, and 5 mM MgSO$_4$). Column was placed inside 15ml falcon tube and centrifuged with 2000 rpm for 4 min at 4°C using Sorvall RT6000 refrigerated centrifuge and repeated same step until the column was packed with Sephadex G 25. Once column was packed 150 µl of buffer (appropriate to the protein) was added and centrifuged with same condition at least 3 times. Column was ready once 150 µl of buffer was coming out as flow through. Autoclaved 1.5ml microtube was used to collect the desalted protein sample.
CHAPTER 3

RESULTS

Section I: Biochemical Characterization of SBIP-428

Bioinformatics Analysis of SBIP-428

DNA and Corresponding Amino Acid Sequences of SBIP-428. In a yeast 2 hybrid (Y2H) screening using SABP2 as a bait, several positive interactors were detected. SBIP-428 was one of the SABP2 interactors. SBIP-428 was then sequenced and analyzed through BLAST search using Sol genomics network database hosting gene sequences of solanaceae plants (Figure 3). SBIP-428 showed homology to a transcriptional regulator Sir2 family protein.

ATGGTTCCTTATTGGATCCCCCTAGCAAGATGTGGACAGTTTGTATGAAATTCTTTTGACAGGATTACCAAGCT
TGTTGATTGGAGCGAGATGGAGACAGAGGAGTGGAAATTCTTTTGACAGGATTACCAAGCT
CTGGTTTAAACAAATTACCACTACGGGACCTTTTTCAGATCATTGATCTTGAGGTCTGAGTATTGGAAGTTAT
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AAGGGATGTGATGCCTTCCTTCTTTACATCTTCTAGGCGAGTCGCAATCGTCAAACCTGAGCAGTTGGGCTTTTGTATGAAAGC

Figure 3: SBIP-428 DNA Sequence Obtained From Y2H
The sequenced DNA was translated using ExPASy Translate tool (http://web.expasy.org/translate/) to obtain corresponding amino acid sequence (Figure 4).

MVPYSDDPSMKDVSILYEFFDRSTKLVLVTAGMMGSTEGSFYSPNGAYSTGFKPITHQEFIRLSVKKARRBYWARSYAGWRRTAQPSTGHIALSSLKAGHISFMITQNVDLHHRAGSSPLELHGTIVVIACTNCGFPLPRELFQDQVKAHNPKWAASVELDAYSRSDESFGKQRPDPGIDIEIDEKFWEDDFYPDRCQGVLPDVVFNGDNVFKARADVAMEAAKGCDASFLVLGSSMNTMSAFRLKAAHEAGAATAIVNIGVTRADDLVPLKINARVGEILPRLLNVGSLIPAL

Figure 4: Corresponding Amino Acid Sequence of SBIP-428 Translated Using ExPASy Bioinformatics Tool.

ExPASy ProtParam tool website was used to predict some other features of SBIP-428. Based on the analysis, the estimated molecular weight of SBIP-428 is 33470.0 Daltons and theoretical pI of 6.25.

Putative Conserved Domains of SBIP-428. Obtained amino acid sequence was then analyzed in “NCBI Standard Protein BLAST” to identify the putative conserved domains using “Non-redundant protein sequence” database. The BLAST results showed that it has a similarity with SIR2 superfamily and specific hits to SIRT4 NAD-dependent deacetylase (Figure 5). It also predicted substrate binding site, NAD⁺ binding site, and Zn binding site. Internal residues involved in substrate binding site are alanine (101, 227, and 258), arginine (223), aspartic acid (225), valine (226), methionine (228), glutamic acid (229), cysteine (135), histidine (256), and glutamic acid (257). Internal residues involved in NAD⁺ binding site are phenylalanine (20), arginine (22), serine (23), threonine (30), glutamine (31), arginine (81), glutamic acid (81), lysine (82), glycine (84), cysteine (135), phenylalanine (214), lysine (279), glutamic acid (257), and isoleucine (280). Internal residues for Zn binding site are proline (143), leucine (146), aspartic acid (211), and phenylalanine (214).
Figure 5: Protein BLAST of SBIP-428. Picture shows the substrate, NAD⁺, and Zn binding sites of SBIP-428 and the putative superfamily protein.

Sequence Alignment with Sirtuin 2 *Arabidopsis thaliana* (AT5G09230) and Human SIR2. ExPASy Bioinformatics Resource Portal ClustalW (http://embnet.vital-it.ch/software/ClustalW.html) sequence alignment tool was used to determine the similarities between SBIP-428, SIR2 (*Homo sapiens*) and *Arabidopsis thaliana* SRT2 (AtSRT2). Sequence alignment showed 77% identity with *Arabidopsis thaliana* (locus AT5G09230) and only 29% with human SIR2 (Figure 6).
Figure 6: Sequence Alignment of SBIP-428, AtSRT2, and Human SIR2. Star at the bottom of alignment shows the identity; dot shows conserved substitution; colon shows semiconserved substitution between 2 proteins; and the gaps showing the dissimilarities between them.

Protein Model of SBIP-428. Putative protein model has been created by using the (PS)$^2$: Protein Structure Prediction Server (http://ps2.life.nctu.edu.tw/). Protein models were created to analyze the structural similarity between SBIP-428, AtSRT2, and crystalized mammalian SIR2 deacetylase. Human SIR2 was used as template to generate SBIP-428 and AtSRT2 protein models. SIR2 deacetylase crystalized structure was obtained from Protein Data Base (PDB) (Yamagata et al. 2014). Comparison between SBIP-428 and AtSRT2 reveals that there are same number of alpha helices and beta sheets in the 3D structure but folding structure is different from
each other (Figure 7). However, SBIP-428 did not show structural similarities because it has only 29% of identity with the template (Figure 8).

**Figure 7:** Protein Model of SBIP-428 and AtSRT2. Predicted 3D structures were designed by using MODELLER 9.13 software. A. 3D structure of SBIP-428 and B. 3D structure of AtSRT2.
Figure 8: Protein Model of SIR2 Deacetylase. SIR2 deacetylase 3D structure (146346.36) was documented in Protein Data Base (PDB) (Yamagata et al. 2014).

Subcellular Localization Prediction of SBIP-428. There are several prediction sites that were used to determine the putative subcellular localization of SBIP-428. Protein Prowler (http://bioinf.scmb.uq.edu.au/pprowler_webapp_1-2/) and MultiLoc (http://abi.inf.uni-tuebingen.de/Services/MultiLoc/) predicted subcellular localization of SBIP-428 in peroxisome whereas BaCelLo (http://gpcr.biocomp.unibo.it/bacello/pred.htm) predicted in nucleus (Figure 9). Same prediction sites were used to find putative subcellular localization for AtSRT2. Same results came out from the predicted sites (data not shown).
**Figure 9:** Predicted Subcellular Localization of SBIP-428. A. prediction using Protein Prowler; B. prediction using MultiLoc; C. prediction using BaCelLo.

**Expression and Purification of SBIP-428**

**Purification of reSBIP-428 Using Ni-NTA Chromatography Expressed in E. coli.**

Recombinant SBIP-428 (6X his tagged) was expressed in *E. coli* and purified using Ni-NTA affinity chromatography. SDS-PAGE and western blot analysis was performed to confirm the purification of rSBIP-428. Samples were mixed with 2X SDS sample buffer, boiled for 5 min, and centrifuged at 13,000 rpm for 2 min prior to load on gel. SBIP-428 was eluted in fraction #4-11. Most of the other proteins except SBIP-428 did not bind and were removed in wash (lane 3) (Figure 10). Fraction # 3-11 contain highest amount SBIP-428 that was detected by western blot using the monoclonal Anti-polyHistidine antibody (Figure 10B).
Figure 10: Purification of SBIP-428 Using Ni-NTA Affinity Column. A. SDS-PAGE gel electrophoresis with eluted fractions (F-3 to F-12) and B. Western-blot analysis of purified rSBIP-428. LMW (low molecular weight) protein marker, input, and wash are presented in first 3 lanes in both pictures. Arrow shows the position of recombinant SBIP-428.

Purification of SBIP-428 Using Mono-Q Ion Exchange Chromatography. Ni-NTA purified fractions (#4-11) containing SBIP-428 were pooled and further purified on a Mono Q column. In Mono Q column most proteins eluted as a sharp peak which was collected as fractions #5-18 (Figure 11). The eluted fractions were analyzed by SDS-PAGE and Western blot. SBIP-428 was detected in the fraction #13-16 (Figure 12B).
Figure 11: Chromatography Profile of Protein in Mono Q Column. (Blue Line) absorbance of protein at 280nm; (Blue Line) salt conductivity; (brown Line) collected fraction (0.5ml).

Fractions #7-16 were used for gel and Western blot analysis.

Figure 12: Purification of SBIP-428 in Mono Q Column. Picture A is protein profile of Mono Q eluted fraction #7-14 in SDS-PAGE (coomassie stain); (LMW) low molecular weight marker. (B) SBIP-428 was detected in fractions #13-16 by Monoclonal Anti-PolyHistidine antibody.
Dialysis of Proteins in Deacetylase Buffer. Mono Q purified SBIP-428 fractions were dialyzed against deacetylase enzyme buffer (50 mM Hepes pH 7.6, 350 mM NaCl, and 20% glycerol). Purified SBIP-428 was pooled together and dialyzed against 500ml deacetylase buffer with slow stirring at 4°C. Buffer was changed every 8 hours for a total of 3 changes. Dialyzed protein was analyzed by SDS-PAGE and detected using the monoclonal anti-polyHistidine antibodies (Figure 13). Dialyzed proteins were aliquoted stored at -80°C until further use.

Figure 13: Dialyzed SBIP-428 After Mono-Q Purification. A. 12.5% SDS-PAGE (Coomassie stained) gel picture; B. Western blot analysis with Monoclonal Anti-PolyHistidine antibody. Arrow shows the position of recombinantSBIP-428.
Deacetylase Enzyme Activity of SBIP-428

Deacetylase Activity Using Total Leaf Protein as Substrate. In order to determine the deacetylase activity of SBIP-428, mono-Q purified SBIP-428 was used and total proteins from the leaves of tobacco plant (XNN) were used as the substrate. Western blot was performed to determine the deacetylase activity of SBIP-428. Acetylated Lysine Antibody was used to detect the deacetylation of acetylated tobacco proteins by SBIP-428. Lane #9 was used as positive control with only substrate proteins and lane #7 was used as negative control with substrate proteins and buffer. Lane’s #1, #3, and #5 contained both enzyme and substrate proteins; #2, #4, and #6 contained only SBIP-428 and the buffer. There was no significant difference between the enzyme treated samples (#1, 3, and 5) and buffer treated controls #7) (Figure 14B). Experiment was performed at least 4 times with similar results.
Figure 14: Deacetylase Activity Assay of SBIP-428; A. Ponceau stained blot, and B. Western blot analysis using Acetylated Lysine Antibody. Pre-stained Marker (PSM) was used as marker proteins. Combination and concentration of substrate (Protein), enzyme (SBIP-428), and buffer are presented at the top.
Deacetylase Enzyme Assay Using SIRT Glo™ Assay Kit. Mono-Q purified SBIP-428 (1 µg, 2 µg, 4 µg, and 8 µg) was used to determine the deacetylase activity using SIRT Glo™ assay kit (Promega). SBIP-428 was expressed, purified, and quantified for this experiment. Deacetylase activity was determined as described in the manufacturer’s protocol. An artificial lysine-acetylated peptide of p53 was used as substrate in luminescence based assay. Free aminoluciferin was measured by Ultra-Glo firefly luciferase reaction that produce a persistent, stable emission of light. SBIP-428 was able to deacetylate the acetylated p53 peptide in linear concentration-dependent manner that indicates that SBIP-428 can act as SIRTUIN-type deacetylase (Figure 15). SIRT Glo™ substrate solution without SBIP-428 was used as negative control in this experiment (Table 6; shown as blank). Luminescence reading was detected using plate reader (BioTek). Raw data and statistical analysis shown in Table 5.

Table 5: Luminescence Reading of SBIP-428 Deacetylase Activity and Statistical Analysis

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Figure 15: Determination of Linear Range of SBIP-428 Enzyme Activity. Recombinant SBIP-428 was serially diluted in 100 µl of SIRT-Glo™ buffer and equal volume of SIRT-Glo™ solution reagent was added in 96-well plate; luminescence signal was detected after 2 hr at room temperature.

Deacetylase Enzyme Assay Using HDAC GloTM Assay Kit. Mono-Q purified SBIP-428 (1 µg, 2 µg, 4 µg, and 8 µg) was used to determine the deacetylase activity using HDAC Glo™ assay kit (Promega). Artificial histone 4 peptide was used as substrate. Free aminoluciferin was measured by Ultra-Glo firefly luciferase reaction that produce a persistent, stable emission of light. SBIP-428 was not able to deacetylate histone 4 peptide in linear concentration-dependent manner that indicates SBIP-428 is not a histone deacetylase (Figure 16). HDAC Glo™ substrate
solution without SBIP-428 was used as negative control in this experiment (Table 6; shown as blank). Luminescence reading was detected using plate reader (BioTek). Raw data and statistical analysis are shown in Table 6

Table 6: Luminescence Reading of SBIP-428 Deacetylase Activity and Statistical Analysis

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Figure 16: Determination of Linear Range of SBIP-428 Deacetylase Activity. Recombinant SBIP-428 was serially diluted in 100 µl of SIRT-Glo™ buffer and equal volume of HDAC-Glo™ solution reagent was added in 96-well plate; luminescence signal was detected after 2 hr at room temperature.

Lysine Acetylation of SBIP-428

Bioinformatics Analysis of SBIP-428 Acetylation on Internal Lysine. Prediction of Acetylation on Internal Lysine (PAIL) ([http://bdmpail.biocuckoo.org/](http://bdmpail.biocuckoo.org/)) was used to predict the putative internal lysine in SBIP-428 that could potentially to get acetylated. Analysis showed that there are 8 lysine residues that could get acetylated (Table 7).
Table 7: Predicted Internal Lysine Residues in SBIP-428

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**Acetylation Detection in SBIP-428.** In order to determine the lysine acetylation of SBIP-428, mono Q purified SBIP-428 was used. Purification of SBIP-428 was determined by western blot analysis using anti-his antibodies (Figure 17). Acetylated bovine serum albumin (BSA) (Fisher Scientific) and albumin (Sigma) were used as positive control and negative control, respectively for this experiment. To determine the lysine acetylation of SBIP-428, Western blot was performed using Acetylated Lysine Antibody. SBIP-428 resulted acetylation in lysine compared with mono-Q purified SBIP-428 (Figure 18).
Figure 17: Western Blot Analysis of Mono Q Purified SBIP-428. A. SDS-PAGE gel electrophoresis picture (arrow indicates SBIP-428) and B. Western blot picture using Anti-Histidine antibody; Lane number 1 in the pictures represent SBIP-428 in deacetylase buffer and lane number 2 with SBIP-428 in Bicine buffer.
Figure 18: Acetylation of SBIP-428. Acetylated bovine serum albumin (BSA) and Albumin (Al) were loaded as positive and negative control, respectively. A SDS-PAGE gel picture (arrow indicates SBIP-428), B. 2 minutes exposure of film, and C. 15 minutes exposure of film in Western blot; Lane number 1 in the pictures represent SBIP-428 in deacetylase buffer and lane number 2 with SBIP-428 in Bicine buffer. Western blot was performed using Acetyl-Lysine Antibody to detect lysine acetylation.
Section II: Acetylation of Internal Lysine Residues of SABP2

SABP2 Acetylation

To determine the acetylation of SABP2, purified SABP2 expressed in *E. coli*, partially purified native SABP2 from tobacco leaves, and native SABP2 from tobacco leaves upon pathogen infection were used.

**Bioinformatics Analysis of SABP2 Acetylation on Internal Lysine.**  
Prediction of Acetylation on Internal Lysine (PAIL) ([http://bdmpail.biocuckoo.org/](http://bdmpail.biocuckoo.org/)) was used to predict the internal lysine in SABP2. Prediction result showed that there are 8 positions where lysine residues have chance to get acetylated (Table 8).

Table 8: Predicted Internal Lysine Residues in SABP2

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Acetylation of Recombinant SABP2. To determine the acetylation of recombinant SABP2, purified SABP2 was used from *E. coli*. Albumin was used as a negative control, whereas acetylated BSA was used as positive control. Recombinant SABP2 (5.25, 7, and 10 µg); egg albumin (10 and 25 µg); and acetylated BSA (0.02 µg) were used for this experiment (Figure 19A). Western blot was performed to detect the acetylated proteins. Recombinant SABP2 did not show lysine acetylation even after extended (5 min) exposure in Western blot (Figure 19B). After prolonged (30 min) exposure, nonspecific background reaction to anti-acetyl lysine antibodies were visualized (Figure 19C).
Figure 19: Acetylation of Recombinant SABP2. A. Ponceau stained of blot; B. Western blot analysis using Acetylated Lysine Antibody at 5min; C. Western blot analysis using Acetylated Lysine Antibody at 30 min.
Acetylation of Native SABP2. Native SABP2 was purified using ammonium sulfate precipitation protocol. 50-75% precipitated proteins were used to determine the acetylation of native SABP2. Western blot was performed using Acetylated Lysine Antibody to detect the acetylation of lysine residue in the proteins (Figure 20C). Proteins were extracted, desalted, quantified, and then subjected to Western blot to detect the acetylation of SABP2 (Figure 20A). Partially purified native SABP2 did not show any internal lysine acetylation (Figure 20C).
**Figure 20:** Acetylation of Native SABP2. A. Coomassie blue stained membrane; B. Western blot analysis using polyclonal SABP2 antibody; C. Western blot analysis using Acetylated Lysine antibody.

**Acetylation of Native SABP2 upon TMV Infection.** *Nicotiana tabacum* cv. Xanthi nc (XNN) tobacco plants were infected with TMV to determine the acetylation of SABP2. Total
soluble protein was extracted and fractionated using ammonium sulfate precipitation. Ammonium sulfate (50-75%) precipitated protein was used to examine SABP2 acetylation (Figure 21). There was no detection of SABP2 acetylation upon TMV infection (Figure 22). 

*PR1* gene expression was analyzed to determine successful pathogen infection (Figure 22).
Figure 21: SABP2 Acetylation upon TMV Infection. A. and B. ponceau stained pictures; C. Western blot analysis using polyclonal SABP2 antibody; D. Western blot analysis using Acetylated Lysine Antibody.

Figure 22: RT-PCR Amplification of PRI Gene. Agarose gel represents expression of PRI at 35 cycle of PCR amplification. DNA ladder (100 bp) was used on the left most lane.

Section III: SBIP-428 in SA Mediated Defense Mechanism

Effect of SA on the Deacetylase Activity in Tobacco Transgenic Plants

To determine the effect of SA on deacetylase activity in plants, tobacco C3, 1-2 (SABP2-silenced), NahG, and Arabidopsis Col-0 were used (Figure 23). Transgenic tobacco 1-2 and NahG plants are accumulate lower levels of SA, whereas C3 and At Col-0 plants were used as control plants. Both NahG and 1-2 tobacco total leaf proteins showed higher levels of lysine acetylation compared with At Col-0 and C3 plant proteins, which implies that deacetylase enzyme as less active in NahG and 1-2 plants (Figure 23C).
Figure 23: Effect of SA on Lysine Acetylated Protein Profile. (A) SDS-PAGE picture of total proteins from tobacco C3, NahG, 1-2, and Arabidopsis Col-0 plants; (B) Ponceau stained blot; (C) Western blot analysis using Acetylated Lysine Antibody.

Effect of SA in Expression of SBIP-428

XNN, NahG, C3, and 1-2 tobacco plants were used for SBIP-428 expression analysis. NahG and 1-2 plants were used to determine the involvement of SA and SABP2 in SBIP-428 expression. XNN and C3 plants were used as wild type and control plats, respectively. EF1α was used as loading control for this experiment, whereas SABP2 gene expression was analyzed to
determine the $SABP2$ silencing in 1-2 tobacco plants (Figure 24). $SBIP$-428 expression was upregulated in NahG plant compared with XNN and C3 plants (Figure 24).

Figure 24: $SBIP$-428 Gene Expression in XNN, NahG, C3, and 1-2 Plants. $SBIP$-428 gene expression after 35 cycles of PCR amplification; $EF1\alpha$ was as loading control; $SABP2$ gene expression in C3 and 1-2 plants as confirmation of $SABP2$ silencing in 1-2 plant.

$SBIP$-428 Expression upon TMV Infection

Six-week-old tobacco plant $Nicotiana tabacum$ cv. Xanthi nc (NN) were infected with TMV to examine the involvement of $SBIP$-428 in plant defense. $PRI$ gene expression was analyzed to determine successful pathogen infection and $EF1\alpha$ gene was analyzed as loading control (Figure 25). $SBIP$-428 gene expression was down regulated at 48 hpi compared with 0, 24, and 78 hpi (Figure 25).
Figure 25: SBIP-428 Gene Expression in XNN Plant upon TMV Infection. SBIP-428 and SABP2 gene expression after 35 cycles of PCR amplification at 0, 24, 48, and 78hpi; EF1-α as loading control; PR1 as confirmation of pathogen infection.

Complementation Assay

Seeds for 6 T-DNA insertion lines of Arabidopsis thaliana SIR2 mutant were obtained from TAIR (arabidopsis.org) to evaluate the role of deacetylase in SA mediate defense signaling. T-DNA insertion lines were SALK_131994C, SALK_149295C, SALK_035541, SALK_139443, CS877409, and CS370961. SALK_139443 and CS370961 mutant lines are histone deacetylase mutant; SALK_131994C, SALK_149295C, SALK_035541, and CS877409 mutant lines are SIR2 deacetylase mutant.
**T-DNA Insertion Confirmation.** All T-DNA insertion lines were analyzed for T-DNA insertion. Col-0 plants were used as control because parent line of all mutants was Col-0. For Col-0, RP and LP primer pair and for all the mutants, LB and RP primer pair was used to examine the T-DNA insertion in the genomic DNA. The expected size of PCR amplified product for mutant lines was 410-710 bp and for Col-0 was 900-1100 bp. All the mutant lines showed the T-DNA insertion (Figures 26 and 27).

![Ladder diagram](image)

**Figure 26:** T-DNA insertion Confirmation Analysis of SRT2 Mutation in *Arabidopsis thaliana.* LB and RP set of primer was used for *Arabidopsis SRT2* mutants and RP and LP set of primer was used for *A. thaliana* Col-0.
**Figure 27:** T-DNA Insertion Confirmation Analysis of *Arabidopsis SRT2* Mutant. LB and RP set of primer was used for *Arabidopsis SRT2* mutants and RP and LP set of primer was used for *A. thaliana* Col-0.

**SRT2 Expression in *Arabidopsis* Mutants.** *SRT2* mRNA expression was analyzed in all *Arabidopsis* mutant along with Col-0. Total RNA was extracted and cDNA was synthesized for RT-PCR amplification of *SRT2* and *Histone Deacetylase. EF1α* was used as loading control for this experiment (Figure 28). Out of 6 seed lines only SALK_131994C line showed altered *SRT2* gene expression (Figure 29A).
Figure 28: RT-PCR Amplification of *EF1α* Gene in *Arabidopsis*. PCR amplification (35 cycles) was used.

Figure 29: RT-PCR Amplification of *SRT2* and *Histone Deacetylase* Gene in *Arabidopsis* Mutants. A. *SRT2* gene mutation analysis and B. *Histone deacetylase* gene mutation analysis. Col-0 was used as positive control for both mutation analysis.

**Confirmation of SRT2 Mutant**. To confirm the *SIRT2* mutation in SALK_131994C seed line, total RNA was extracted from mature leaves, cDNA was synthesized, and *AtSRT2* gene analyzed by RT-PCR. SALK_131994C seed line showed *SRT2* mutation (Figure 31). *EF1α* gene was used as loading control (Figure 30).
Figure 30: AtSRT2 Gene Analysis in SRT2 Mutant. AtSRT2 gene expression after 30 cycle of PCR amplification; EF1α gene was analyzed as loading control.

Growth Phenotype of Arabidopsis SRT2 Mutant (SALK_131994C)

To determine the morphological difference between Arabidopsis SRT2 mutant (SALK_131994C) and parent Arabidopsis thaliana Col-0, 50 seeds of each mutant and control were grown in autoclaved soil. Sown seeds were kept in dark and cold condition for 3 days and then transferred to the growth chamber with 700 to 800 lux light with 16h day light period. Germination rate was determined after 7 days of sowing seeds. Number of leaves on each plant was counted and length of leaves were measured after 21 days of germination. Number of flowers and length of flowers bolts were measured between 30 to 35 days. For seed germination, 50 seeds were sown for each mutant and control as described above. After 7 days, number of germinated seeds was counted and data analyzed. Germination rate of Arabidopsis SRT2 mutant
was 98% and Col-0 was 96% (Figure 32A). There was no significant difference in germination rate between SIR2 mutant and Col-0 seeds.

Figure 31: Growth Phenotype of Arabidopsis SRT2 Mutant. A. Grown seedling of SRT2 mutant; B. Plants growing in growth chamber; C. Measurement of flower bolts.

To examine the leaf morphology of Arabidopsis SRT2 mutant, number of leaves was counted as well as length of leaves were measured. Leaf morphology was also examined visually. All 50 plants leaves were measured and counted. The average leaves number of Arabidopsis SRT2 mutant was 15.8 and Col-0 was 15.3 (Figure 32B). There was no significant
difference in leaves number between Col-0 and SRT2 mutant. Lengths of leaves were measured using roller scale. The average length of the Arabidopsis SRT2 mutant leaves was 3.45cm and Col-0 was 3.56cm (Figure 32C). Length of leaves showed no significant difference between SRT2 mutant and Col-0.

Length of the flower bolts was measured after 35 days of germination (Figure 32D). There were no visually distinguishable difference in floral morphology as well as as determined by statistical analysis. Overall growth phenotype study showed no significant difference between Col-0 and SIR2 mutant.

Figure 32: Growth Phenotype of Arabidopsis thaliana Col-0 and SRT2 Mutant. A. Germination rate; B. Number of leaves per seedlings; C. length of each leaf; D. Height of flower bolt.
**Pathogen Growth Assay**

*Arabidopsis thaliana* wild type Col-0 and SRT2 mutant seeds were grown in 1/2 strength MS media (Figure 33). Twenty days old plants were used for to pathogen growth assay. Flooding inoculation technique was used to inoculate plants with pathogen. *Pst* DC3000 and *Pst* DC3000 AvrRpt2 bacterial strains were used for plant inoculations at concentration of $1 \times 10^8$ cfu/ml and $5 \times 10^6$ cfu/ml. Both Col-0 and SRT2 mutant plants inoculated with $1 \times 10^8$ cfu/ml started to show chlorosis symptoms at 2 dpi and died at 4 dpi (Figure 34). With concentration of $5 \times 10^6$ cfu/ml plants started to show chlorosis at 4 dpi (Figure 34). For control, plants were treated with water. Bacterial growth was observed at 4 dpi and colony forming units (cfu/ml) were counted after 2 and 3 days of plating. In Col-0, bacterial population was $6.0 \times 10^7$ and $7.1 \times 10^7$ cfu/ml when the plans were treated with *Pst* DC3000 and *Pst* DC3000 AvrRpt2, respectively (Table 9). Whereas in SRT2 mutant bacterial population was $2.3 \times 10^7$ and $1.18 \times 10^7$ cfu/ml when plants were inoculated with *Pst* DC3000 and *Pst* DC3000 AvrRpt2, respectively (Table 9). Bacterial growth was more than 3 times higher in Col-0 compared with SRT2 mutant in plants were treated with *Pst* DC3000 (Figure 35). The bacterial growth was 7 times higher in plants inoculated with *Pst* DC3000 AvrRpt2 (Figure 36). A comparison growth of *Pst* DC3000 and *Pst* DC3000 AvrRpt2 bacterial growth in Col-0 and SRT2 mutant is shown in Figure 37.
Figure 33: Growth Phenotype of *Arabidopsis SRT2* Mutant in 1/2 MS media. A-C. *Arabidopsis thaliana* Col-0 growth at 7, 14, and 21 days; D-F. *Arabidopsis thaliana SRT2* mutant growth phenotype at 7, 14, and 21 days.
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**Figure 34** (continued on the next page)
Figure 34: Disease Phenotype of *Arabidopsis thaliana* at 2 dpi and 4 dpi. A-D. H₂O treated *Arabidopsis* Col-0 and SRT2 mutant; E-H. *Pst* DC3000D 10⁸ cfu/ml inoculated *Arabidopsis* Col-0 and SRT2 mutant; I-L. *Pst* DC3000D *AvrRpt2* 10⁸ cfu/ml inoculated *Arabidopsis* Col-0 and SRT2 mutant; M-P. *Pst* DC3000D 5 X 10⁶ cfu/ml inoculated *Arabidopsis* Col-0 and SRT2 mutant; Q-T. *Pst* DC3000D *AvrRpt2* 5 X 10⁶ cfu/ml inoculated *Arabidopsis* Col-0 and SRT2 mutant.

Table 9: Bacterial Population (*Pst* DC3000 and *Pst* DC3000 *AvrRpt2*) in Col-0 and SRT2 Mutant at 4 dpi

<table>
<thead>
<tr>
<th></th>
<th><em>Pst</em> DC3000 (cfu/mg)</th>
<th><em>Pst</em> DC3000 <em>AvrRpt2</em> (cfu/mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Col-0</td>
<td>6.0 x 10⁷</td>
<td>7.1 x 10⁷</td>
</tr>
<tr>
<td>SRT2 mutant</td>
<td>2.3 x 10⁷</td>
<td>1.18 x 10⁷</td>
</tr>
</tbody>
</table>
Figure 35: Bacterial Population of \textit{Pst} DC3000 in \textit{A. thaliana} (Col-0) and \textit{A. thaliana} SRT2 Mutant flood-inoculated with a concentration of $5 \times 10^6$ cfu of bacterial suspension. Bacterial populations were quantified at 4 dpi.

Figure 36: Bacterial population of \textit{Pst} DC3000 \textit{AvrRpt2} in \textit{A. thaliana} (Col-0) and \textit{A. thaliana} SRT2 mutant flood-inoculated with $5 \times 10^6$ cfu of bacterial suspension. Bacterial populations were quantified at 4 dpi.
Figure 37: A Comparison Graph between *Pst* DC3000 and *Pst* DC3000 AvrRpt2 Growth in *Arabidopsis* Col-0 and *Arabidopsis* SRT2 Mutant at 4 dpi.
CHAPTER 4

DISCUSSION

Understanding the SA signaling pathway in the development of SAR and the involvement of SBIP-428 are the central foci of this project. SABP2 helps to convert lipid soluble MeSA to SA in order to increase the plant's resistance (Kumar and Klessig 2003). SABP2 likely helps to increase the cytoplasmic SA levels that results in changes in the redox potential of the cytoplasm (Tripathi et al. 2010). However, the protein interaction cascade is poorly understood in the SABP2-mediated SA signaling pathway. To understand the interaction cascade in this pathway, a yeast 2 hybrid (Y2H) screening was performed to identify the SABP2 interacting proteins. Several proteins were identified as Salicylic Acid Binding Protein 2 Interacting Proteins (SBIPs). SBIP-428 is one of the interacting proteins. Characterization of SBIP-428 and determining its role in the SABP2 dependent SA signaling pathway has been the main focus of this study. Bioinformatics analysis of SBIP-428 suggested that it belongs to the SIR2 super family of NAD\(^+\) dependent deacetylases. These group of proteins regulate the acetylation status of proteins. Based on bioinformatics, our first attempt was to determine if SBIP-428 is a true NAD\(^+\) dependent deacetylase. Interaction of SABP2 with SBIP-428, a deacetylase raises several question: why does SBIP-428 interact with SABP2? Is SABP2 itself acetylated, which may regulate the enzyme activity? Does acetylation/deacetylation play any significant role in the SABP2 mediated SA signaling defense pathway? How does this interaction between SABP2 and SBIP-428 affect the plant defense pathway? To answer some of these questions, we hypothesized that SABP2 is acetylated and is regulated by the deacetylase activity of SBIP-428, and that deacetylase activity of SBIP-428 has an effect on SA mediated defense mechanism.
Biochemical Characterization of SBIP-428

Bioinformatics analysis, protein BLASTS analysis with nonredundant protein sequence, and Protein Data Base (PDB) analysis have shown that SBIP-428 is a potential NAD\(^+\) dependent SIR2 deacetylase. Translated SBIP-428 protein sequence shows 93% identity with predicted NAD-dependent protein deacetylase SRT2-like (\textit{Solanum lycopersicum}) and 77% identity with sirtuin 2 \textit{Arabidopsis thaliana} (AtSRT2) (AT5G09230). However, BLAST analysis with PDB showed 29% identity with human Sir2 protein that is bound to an acetylated p53 peptide (Avalos et al. 2002).

Few other bioinformatics tools were used to predict the potential localization and signal peptide of SBIP-428. According to “MultiLoc” and “Protein Prowler”, SBIP-428 is potentially localized in peroxisomes. And based on the information obtained from the BaCelLo website, SBIP-428 could potentially be localized in the nucleus (Figure 9). AtSRT2, a negative regulator of SA pathway, was reported in nucleus. However, most recently same AtSRT2 was reported in inner mitochondrial membrane involved in energy metabolism and transportation (Konig et al. 2014). SABP2 is likely localized on the chloroplast outer envelope membranes (Fai and Kumar, unpublished). SBIP-428 was identified as an interactor of SABP2 in yeast 2 hybrid and a physical interaction was determined by pull down assay using a GST-tagged SBIP-428 (Zhao and Kumar unpublished). With respect to SABP2 localization, SBIP-428 may be localized in the nucleus or peroxisomes and it transport to chloroplast outer envelope to interact with and regulate SABP2 activity when is needed. There are reports showed that sirtuin deacetylases can be localized in various subcellular compartments. SIR2 super family proteins can be localized either in nucleus, cytoplasm, or mitochondria (Argmann and Auwerx 2006; Bjoern et al. 2006; Ahuja et al. 2007; Du et al. 2011; Kim et al. 2011; Barber et al. 2012; and Sundaresan et al.)
SIRT3 is localized in nucleus and mitochondria (Bunkenborg et al. 2006; Hirschey et al. 2010). SIRT4 that is found in mitochondria in mammals (Ahuja et al. 2007).

Enzyme assay for recombinant purified SBIP-428 was conducted by following the modified deacetylase enzyme activity protocol from Finkemier et al. (2011). Instead of using a specific substrate, total naturally acetylated proteins from tobacco leaf (XNN) was used as a substrate for the deacetylase activity of SBIP-428. There was no significant difference between the deacetylase activity of SBIP-428 and the negative control of buffer (Figure 14), suggesting presence of significant deacetylase activity in tobacco leaves (Figure 14B). Therefore, it was not possible to determine the enzyme activity of SBIP-428 using total acetylated leaf proteins of tobacco.

SIRT-Glo™ and HDAC-Glo™ assay kits (Promega) were used to test the deacetylase activity of SBIP-428. Lysine-acetylated peptide of p53 as substrate in SIRT-Glo™ assay kit whereas histone 4 peptide was used as substrate in HDAC-Glo™ assay kit. Mono-Q purified SBIP-428 was used to determine the enzyme activity. Results showed that SBIP-428 was able to deacetylate artificial p53 peptide but not histone 4 peptide in linear concentration-dependent manner (Figures 15 and 16). Ability of p53 peptide deacetylation suggests that SBIP-428 is a sirtuin deacetylase. Sirtuin deacetylase are involved in aging, glucose homeostasis, fatty acid oxidation, control of gene expression, etc. (Argmann and Auwerx 2006; Hirschey et al. 2009; Zhong et al. 2010). Most recently report suggests that Sirtuin deacetylase in Arabidopsis plays a role in energy metabolism and metabolic transportation (Konig et al. 2014). Role of sirtuin deacetylases in plant defense yet to be discovered.
Lysine acetylation is a common posttranslational modification that is readily reversible with deacetylation (Finkemier et al. 2011). In certain instance of biological equilibrium, lysine acetylation and deacetylation can occur on the same protein. So far, there are no reports that showed deacetylase enzyme itself is regulated by acetylation/deacetylation. Bioinformatics analysis was performed to predict the potential lysine residues in SBIP-428 (Table 7). Bioinformatics tool for lysine acetylation prediction sites and experimental results were both positive. More than one prediction site was used for lysine acetylation prediction and all of them showed SBIP-428 has potential lysine residues for acetylation. One analysis showed 9 potential acetylation sites and another site showed only 1 site. To biochemically determine if SBIP-428 was itself acetylated, mono Q purified SBIP-428 was subjected to western blot analysis using anti acetyl-lysine antibodies. Results show that SBIP-428 is acetylation (Figure 18). Acetylation of SBIP-428 (sirtuin deacetylase) lead us to believe that a posttranslational modification may be needed for its function and controlled by other regulatory factors. This is similar to another common posttranslational modifications mediated by the MAPK cascades in which MAPK is phosphorylated by MAPKK and MAPKK is by MAPKKK.

Acetylation of SABP2

Interaction of SBIP-428 (a deacetylase) with SABP2 raised the possibility of SABP2 itself being acetylated which could potentially be deacetylated by SBIP-428. Bioinformatics tool for lysine acetylation prediction sites show that SABP2 has potential lysine residues for acetylation (Table 8). Recombinant SABP2 and native SABP2 (partially purified from uninfected or TMV infected tobacco leaves) was used to determine acetylation of SABP2. However, SABP2 (recombinant or native) did not show any acetylation (Figures 19, 20, and 21). Recombinant SABP2 was expressed and purified from *E. coli*. Because acetylation and
deacetylation is a readily reversible reaction, it might be possible that SABP2 becomes acetylated under certain conditions and is deacetylated under another conditions.

**SBIP-428 in Plant Defense**

Results showed SA has little impact on deacetylase activity in tobacco transgenic plants (Figure 23). Plants known to accumulate or synthesize less SA showed fewer deacetylation of naturally acetylated proteins. Compared with total leaf proteins from C3 tobacco plants (control plant), proteins from NahG and 1-2 plants showed higher lysine acetylation, which suggest that deacetylase enzyme is less active in plants that produce/accumulate less SA. However, proteomics analysis is needed to specify the proteins whose acetylation is affected by SA levels in plants.

The same NahG, 1-2, C3, and XNN plants were used for **SBIP-428** gene expression. NahG transgenic plants always showed higher expression of **SBIP-428** compared with other transgenic and control plants (Figure 24). It suggests that SA and SBIP-428 regulate each other negatively, which is affirmative to the observation of AtSRT2, a homolog of SBIP-428 (Wang et al. 2010). Wang et al. (2010) showed that AtSRT2 negatively regulates the SA defense pathway by suppressing defense related genes as well as the genes related to SA synthesis.

To determine the role of SBIP-428 in host-pathogen interactions, TMV inoculated plants were used. XNN plants were infected with TMV and expression of SBIP-428 was examined at 0, 24, 48, and 78hpi. Primary observation showed that **SBIP-428** expression was down-regulated and **PRI** expression was up-regulated at the 48hpi (Figure 25). These opposite expression results suggest that expression of **PRI** and **SBIP-428** are inversely correlated. Gene expression is closely related to histone deacetylase and acetyltransferase activity (Choudhary et al. 2009). Less
expression of histone deacetylase facilitate the expression of other genes because more
deacetylase activity means more tightly binding of histone to DNA and less genes are allowed to
be expressed. Therefore, inverse expression of SBIP-428 and PRI suggests that less expression
of putative sirtuin deacetylase (SBIP-428) somehow regulates the expression levels of PRI gene.
As the SIR 2 super family deacetylase is diverse in localization, it is possible that SBIP-428
decaylsyl enzyme localized in nucleus and regulates PRI gene expression.

The AtSRT2 mutant (SBIP-428 homolog) was studied to understand the effect of
deacetylase in the SA defense pathway. Primarily, 6 T-DNA insertion mutants were obtained
from TAIR. T-DNA insertion analysis (Figures 26 and 27), AtSIR2 gene expression analysis
confirmed that one of the T-DNA insertion line, SALK_131994C line was a true mutant (Figures
28-30). SALK_131994C was used for growth phenotype and bacterial growth assay analysis.
Growth phenotype studies showed no significant difference in terms of seed germination, leaf
morphology, and flower morphology (Figures 31 and 32).

Wang et al. (2010) have reported the putative deacetylase AtSRT2 function in plant
defense. They reported that AtSRT2 deacetylase negatively regulates the plant defense and the
expression of PAD4, EDS5, SID2, and PRI was down regulated. EDS5 and SID2 are directly
related and PAD4 is indirectly responsible for the production of SA. The suppression of these
genes indicate that AtSRT2 deacetylase activity is inhibitory to the production of SA and plays a
negative role in SA dependent plant defense signaling.

Pathogen growth assay using virulent Pst DC3000 showed the similar effect reported by
Wang et al. (2010). The AtSRT2 mutant plants were more resistant compared to Col-0 as
observed by Wang et al. (2010). Plants started to show chlorotic symptoms at 2dpi and died at
4dpi with $1 \times 10^8$ cfu in both Col-0 and AtSRT2 mutant, which suggest that high concentration of bacterial inoculation did not make any difference in terms of disease resistance (Figure 33). However, at low bacterial inoculation ($5 \times 10^6$ cfu), plants survived till 4 dpi. Chlorosis symptom was observed only in Col-0 plants and bacterial growth was 3-fold higher compared with AtSRT2 mutants (Figure 34).

To investigate the effector-triggered immunity (ETI), Pst DC3000 carrying AvrRpt2 was used. There was no difference compared with Pst DC300 when plants were inoculated with high concentration of $1 \times 10^8$ cfu. Plants showed similar chlorosis symptoms at 2 dpi and died at 4 dpi (Figure 34). But when plants were inoculated with $5 \times 10^6$ cfu, they showed different response. AtSRT2 mutant showed better resistance compared with Col-0. Pst DC3000 AvrRpt2 grew 7 times higher in Col-0 (Figure 36). Results suggests that AtSRT2 mutant showed better resistance against both virulent and avirulent bacterial pathogen. These results demonstrate that AtSRT2 mutant has a better ETI response compared with wild type, which leads us to believe that AtSRT2 is a negative regulator of basal defense.

This research was conducted to characterize the SBIP-428 and to determine its role in SA signaling defense pathway. The results presented in this thesis showed that SBIP-428 is a sirtuin decetylase enzyme. Expression of SBIP-428 revealed that SBIP-428 may have negative correlation with SA. Arabidopsis SRT2 mutant showed no morphological differences in absence of SRT2 gene. Mutant study also revealed that mutant plant has better ETI response and AtSRT2 negatively correlated in basal defense. Absence of SRT2 make plant more resistant. Another interesting finding was to detect the acetylation of SBIP-428.
Future Direction

Characterization of SBIP-428 is just a beginning of this project. To get better understanding the role of SBIP-428 in plant defense, it will be important to generate transgenic tobacco silenced in SBIP-428 expression or to overexpress it. These transgenic tobacco plants could be used to determine the role of SBIP-428 on the host and nonhost defense responses.

Also, we have shown the first time that deacetylase enzyme itself gets acetylated. It will be important to investigate the deacetylase activity of SBIP-428 by altering the potential lysine residues that are acetylated. Changes in acetylation of SBIP-428 may lead to changes in its catalytic function.
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APPENDICES

Appendix A – Abbreviations

SABP2 - Salicylic acid binding protein 2
SBIP-428 – SABP2 Interacting Protein-428
C3 - Control plants (*Nicotiana tabacum* cv Xanthi nc, a local lesion host of Tobacco Mosaic
Virus and contains empty silencing vector.
NahG - Plants expressing salicylate hydroxylase which converts SA to catechol.
1-2 - SABP2 - silenced plants (transgenic *N.t. cv Xanthi nc* in which *SABP2* gene expression is
silenced by RNA interference.
PRRs - Pattern recognition receptors
PAMPs - Pathogen-associated molecular patterns
R protein - Resistance protein
Avr - Avirulence
ICS 1 - Isochorismate synthase 1
BA2H - Benzoic-2-hydroxylase
HR - Hypersensitive response
PCD - Program cell death
SA - Salicylic acid
JA - Jasmonic acid
ET - Ethylene
ISR - Induced systemic resistance
SAR - Systemic acquired resistance
SAMT - Salicylic acid methyl transferase
MeSA - Methyl salicylate

NO - Nitric oxide

SDS PAGE - Sodium dodecyl sulphate-polyacrylamide gel electrophoresis

TMV - Tobacco mosaic virus

PR - Pathogenesis-related

BTH - Benzo-(1, 2, 3)-thiadiazole-7-carbothioic acid S-methyl ester

INA - 2, 6-dichloro-isonicotinic acid

ROI - Reactive oxygen intermediates

NPR1 - Non-expresser of pathogenesis-related protein 1

IPL - Isopyruvate lyase

NFAT - Nuclear factor of activated T-cells 87

cfu - colony forming units

βME - βeta mercaptoethanol

\textit{Pst} - \textit{Pseudomonas syringae}

\textit{EFalpha1} - \textit{Elongation Factor alpha 1}

\textit{PAD4} - \textit{Phytoalexin Deficient 4}

\textit{R}-genes - \textit{Resistance genes}

RLK - Receptor-like kinase

RLP - Receptor-like proteins

TAE - Tris-Acetate EDTA

KBM - King’s B Medium

KDa - Kilo Dalton

OD - Optical Density
UV - Ultra violet

μg - micro gram

μl - micro litre

ml - milli litre

mM - milli Molar
Appendix B – Buffers and Reagents

Protein Extraction Buffer (buffer A) (1L)

Sodium Citrate (7.44g), M.W. = 372.24g/L, final concentration = 20mM

MgSO4 (1.23g). M.W. = 246.48g/L, final concentration = 5mM

EDTA (0.42g), M.W. = 416.20g/L, final concentration = 1mM

Adjust pH to 6.3

Stored at 4°C until use.

Prior to grinding plant tissue add 1ml of β-ME (14.4mM final concentration), 1ml of PMSF (100mM) (0.1mM final concentration), 0.15g of benzamidine HCl (156.61g/mol, final concentration 1mM ) and 15g of 100% PVPP (1.5% wt/vol) to 1L of buffer.

KING’S B Medium

Protease peptone # 3 = 20 g

Potassium phosphate dibasic = 1.50 g

Magnesium sulfate = 1.50 g

Glycerol = 10 ml

Adjust the volume to 1 liter with distilled water

Adjust the pH to 7.0

Agar = 17.50 g (for solid medium)

Autoclave for 30 minutes before use

10 mM Magnesium Chloride

MgCl2 = 0.952 g

Adjust the volume to 1 liter with distilled water

1M Magnesium Sulfate
MgSO4 = 246.48 g

Adjust the volume to 1 liter with distilled water

0.1M Sucrose Solution

Sucrose = 34.2 g

Adjust the volume to 1 liter with distilled water.

Filter sterilize the solution and store at -20°C

0.1% Diethyl Pyrocarbonate Treated Water

Diethyl pyrocarbonate = 0.1 ml

Add to 100 ml distilled water

Incubate for ~12 hours at 37°C

Autoclave for 15 minutes

RIFAMPICIN (14 mg/ml)

Rifampicin (powder) = 0.14 g

Add to 10 ml of Methanol

Add to King’s B media at 25 μg/ml

Bicine buffer (buffer B) (1L)

Bicine (1.63g), M.W. = 163.2g/mol, final concentration = 10Mm

Adjust pH to 8.0 with 1 N NaOH

10x Phosphate Buffer Saline (10x PBS)

Sodium Chloride (76g), M.W. = 58.44g/mol, final concentration = 1.3M

Sodium Phosphate dibasic (10g), M.W. = 141.96g/mol, final concentration = 70mM

Sodium Phosphate monobasic (4.1g), M.W. = 119.96g/mol, final concentration = 30mM

For 1x PBS (1 L), dilute 100mL of 10x PBS in 900mL of water. 88
For 1x PBS (1 L) with 3% Tween 20, dilute 100mL of 10x PBS in 870mL, then add 30mL of tween 20.

Western Blotting Blocking Buffer (100mL)

1x PBS buffer, 100mL

Dry Milk (1g), final concentration = 1%

BSA (3g), final concentration = 3%

4x SDS-PAGE Stacking gel Buffer (500mL)

Tris base (30.28), M.W. = 121.1g/mol, final concentration = 0.5M

Adjust pH to 6.8

Add SDS (0.2g), final concentration = 0.04%

SDS Gel Loading Buffer (2 X):

50mM Tris-HCl (pH 6.8)

100mM DTT

2% (wt/vol) SDS

0.1% (wt/vol) bromphenol blue

10% (vol/vol) glycerol

10x SDS-PAGE Running Buffer (1 L)

Tris base (30g), M.W. = 121.1g/mol

Glycine (144g), M.W. 75.07g/mol

SDS (10g) 89

10x Western Blotting Transfer Buffer (1L)

Tris base (30.3g), M.W. = 121.1g/mol, final concentration = 125mM

Glycine (72.06g), M.W. = 75.07g/mol, final concentration 960mM
For western, 1x transfer buffer is prepared by mixing 100mL of 10x transfer buffer, 100mL of 100% methanol, and 800mL of cold water.

Ammonium Persulfate (20% in 1mL)
Dissolve Ammonium persulfate (20mg) in 1mL of water

2x SDS-PAGE Loading Dye (100mL)
1M Tris-Cl, pH 6.8 (10mL), final concentration = 100mM
SDS (0.4g), final concentration = 0.4%
Glycerol (20mL), final concentration, 20%
Bromophenol blue (0.2g), final concentration = 0.2%
Add 5mL of ME before use.

Ponceau S Stain (100mL)
Ponceau S (0.1g), final concentration = 0.1%
Acetic acid (5mL), final concentration = 5%

MS media with Gamborg’s Vitamins
MS media (4.4g/L)
Myo-Inositol (100mg/L)
Nicotinic Acid (1mg/L)
Pytodoxine. HCl (1mg/ml)
Thiamine. HCl (10mg/ml)
VITA

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